

Practitioner's Docket No. 700953-45394C

PATENT

Preliminary Classification: Proposed Class:

Subclass:

NOTE: "All applicants are requested to include a preliminary classification on newly filed patent applications. The preliminary classification, preferably class and subclass designations, should be identified in the upper righthand corner of the letter of transmittal accompanying the application papers, for example 'Proposed Class 2, subclass 129." M.P.E.P. Section 601, 7th ed.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Box Patent Application Assistant Commissioner for Patents Washington, D.C. 20231

NEW APPLICATION TRANSMITTAL

Transmitted herewith for filing is the patent application of Inventor(s): Jeffrey, SCHLOM; Dennis, PANICALI

CERTIFICATION UNDER 37 C.F.R. SECTIONS 1.8(a) AND 1.10*

(When using Express Mail, the Express Mail label number is mandatory; Express Mail certification is optional.)

I hereby certify that, on the date shown below, this correspondence is being:

		MAILING					
[]	deposited with the United States Postal Service in an envelope addressed to the Assistant Commissioner for Patent						
	Washington, D.C. 20231. 37 C.F.R. Section 1.8(a)	37 C.F.R. Section 1.10*					
[]	with sufficient postage as first class mail.	[x] as "Express Mail Post Office to Address" Mailing Label No(mandator) EL565095244	y) 145				
	TRA	NSMISSION					
[]	transmitted by facsimile to the Patent and Tradem	Patricia survi					
Date: _	October 20 2000	· · · · · · · · · · · · · · · · · · ·	<u></u>				
		Signature					
		Patricia Turner					
		(type or print name of person certifying)					

*WARNING:

Each paper or fee filed by "Express Mail" must have the number of the "Express Mail" mailing label placed thereon prior to mailing. 37 C.F.R. Section 1.10(b).

"Since the filing of correspondence under [Section] 1.10 without the Express Mail mailing label thereon is an oversight that can be avoided by the exercise of reasonable care, requests for waiver of this requirement will not be granted on petition." Notice of Oct. 24, 1996, 60 Fed. Reg. 56,439, at 56,442.

(New Application Transmittal--page 1 of 13)

1.

For (title):GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC ANTIGEN (PSA)

Type of Application

This ne	w applic	cation is for a(n)
	[] [] [] [X] []	Original (nonprovisional) Design Plant Divisional. Continuation. Continuation-in-part (C-I-P).
2.	Benefit	t of Prior U.S. Application(s) (35 U.S.C. Sections 119(e), 120, or 121)
	[X]	The new application being transmitted claims the benefit of prior U.S. application(s). Enclosed are ADDED PAGES FOR NEW APPLICATION TRANSMITTAL WHERE BENEFIT OF PRIOR U.S. APPLICATION(S) CLAIMED.
3.	Papers	Enclosed
	A.	Required for Filing Date under 37 C.F.R. Section 1.53(b) (Regular) or 37 C.F.R. Section 1.153 (Design) Application
		Pages of Specification Pages of Claims Sheets of Drawing
	В.	Other Papers Enclosed 1 Pages of declaration and power of attorney Pages of Abstract Other
4.	Additi	onal Papers Enclosed
	[]	Amendment to claims
		[] Cancel in this applications claimsbefore calculating the filing fee. (At least one original independent claim must be retained for filing
		purposes.) [] Add the claims shown on the attached amendment. (Claims added have been

(New Application Transmittal--page 2 of 13)

5.

is submitted.

will be submitted.

[]

[]

			numbe	ered consecutively following the highest numbered original claims.)
		Inform Form I Citation Declar Submit pertain sequen Author	nation D PTO-14 ons ration of ssion of ning then	mendment disclosure Statement (37 C.F.R. Section 1.98) 49 (PTO/SB/08A and 08B) EBiological Deposit Sequence Listing," computer readable copy and/or amendment reto for biotechnology invention containing nucleotide and/or amino acid of Attorney(s) to Accept and Follow Instructions from Representative nents
5.	Declar	ration o	r Oath	(including power of attorney)
	[X]	Enclos	sed	
		Execu	ted by	
		[X] [] []	joint i	tor(s). representative of inventor(s). 37 C.F.R. Section 1.42 or 1.43. nventor or person showing a proprietary interest on behalf of inventor who do sign or cannot be reached.
			[]	This is the petition required by 37 C.F.R. Section 1.47 and the statement required by 37 C.F.R. Section 1.47 is also attached. See item 13 below for fee.
	[]	Not E	nclosed.	
		[]		cation is made by a person authorized under 37 C.F.R. 1.41 on behalf of almove named inventor(s).
			[]	Showing that the filing is authorized. (not required unless called into question. 37 C.F.R. Section 1.41(d))
6.	Inven	torship	Statem	ent
The in	ventorsl	nip for a	ll the cla	aims in this application are:
	[]	The sa	ame.	or
	[]			An explanation, including the ownership of the various claims at the time ed invention was made,

Language							
[X] []	_						
	[]			at the translation is accurate. 3			
Assignment							
[]	An ass	signment of the invention	on to				
	[]	MENT) ACCOMPA	NYING NEW PATENT A	OR ASSIGNMENT (DOCU- PPLICATION" or [] FORM			
	[]	will follow.					
	_						
Со	untry		Appln. no.	Filed			
Со	untry		Appln. no.	Filed			
Со	untry		Appln. no.	Filed			
which p [] []	is (are	e) attached.					
Fee (Calculati	ion (37 C.F.R. Section	1.16)				
A.	[X]	Regular application					
	Assig [] Certif Co Co Co Fee C	Assignment [] An ass [] Certified Copy Certified copy Country Country Country Country Fee Calculations Fee Calculations	[] Non-English [] The attached translatic C.F.R. Section 1.52(d) Assignment [] An assignment of the invention of the in	[] Non-English [] The attached translation includes a statement that C.F.R. Section 1.52(d). Assignment [] An assignment of the invention to [] is attached. A separate [] "COVER SHEET FOMENT) ACCOMPANYING NEW PATENT ALL PTO 1595 is also attached. [] will follow. Certified Copy Certified copy(ies) of application(s) Country Appln. no. Country Appln. no. Country Appln. no. which priority is claimed [] is (are) attached. [] will follow. Fee Calculation (37 C.F.R. Section 1.16)			

Claims		Number Filed	Basic Fee Allowance	Number Extra	Rate	Basic Fee 37 C.F.R. Section 1.16(a) \$710.00
Total C (37 C.F Section 1.16(c)	.R.		18 - 20 =	X	\$18.00	
-	ndent Cla LR. Sect		4 - 3 =	X	\$78.00	\$78.00
Claim(s	e Depens), if any C.R. Sect	,				
			+	\$260.00		
	[]	Amendment canc Amendment delet Fee for extra clair	ing multiple-depe	endencies is enclose	ed.	
			F	iling Fee Calculation	on	\$788.00
	В.	[] Design a (\$310.0037 C.F		(f)) iling Fee Calculation	on	\$
	C.	[] Plant app (\$480.0037 C.F	R. Section 1.16	(g)) iling Fee Calculation	on	\$
11.	Small I	Entity Statement(s)			
	[]	Statement(s) that (are) attached.	this is a filing by	a small entity und	ler 37 C.F.F	R. Section 1.9 and 1.27 is
	[]		, filed on	d in prior applicatio		which benefit is being

(New Application Transmittal--page 5 of 13)

		35 U.S.	.C. Section	[] [] []	119(e), 120, 121, 365(c),				
		and wh	ich status as a	small ent	ity is still proper and d	esired.			
		[]	A copy of the	e statemer	nt in the prior application	on is includ	led.		
		Filing 1	Fee Calculatio	on (50% of	f A, B or C above)	\$	788.00)	
12.	Reque	st for In	ternational-T	ype Searc	ch (37 C.F.R. Section 1	.104(d))			
	[]	Please nationa	prepare an ir	nternations	al-type search report i erits takes place.	for this ap	plication	at the time	e when
13.	Fee Pa	ayment F	Being Made at	t This Tin	ne				
	[]	Not Enclosed							
		[]	No filing fee (This and to subsequently	the surch	paid at this time. arge required by 37	C.F.R. Se	ction 1.1	16(e) can i	be paid
	[X]	Enclos	sed						
		[X]	Filing fee				\$	788.00	•
		[]	(See attache	C.F.R. Se ed "COVE ENT ACC	t ection 1.21(h)) ER SHEET FOR COMPANYING NEW		\$	<u></u>	_
		[]	be reached	inventors f the inver fused to si	or person	(i))\$			
		[]	specification	n in a nor	plication with a n-English language Sections 1.52(d) and 1.	17(k))\$			

		[]	Processing and retention fee (\$130.00; 37 C.F.R. Sections 1.53(d) and 1.21(l))\$	
		[]	Fee for international-type search report (\$40.00; 37 C.F.R. Section 1.21(e))	\$
			Total Fees Enclosed	\$
14.	Metho	d of Pay	ment of Fees	
	[X]	Check	in the amount of \$	
	[]	_	e Account No in the amount of \$ licate of this transmittal is attached.	•
15.	Autho	rization	to Charge Additional Fees	
	[X]	The C	Commissioner is hereby authorized to charge the for and during the entire pendency of this application to	ollowing additional fees by this Account No. 50-0850
		[X]	37 C.F.R. Section 1.16(a), (f) or (g) (filing fees)	
		[]	37 C.F.R. Section 1.16(b), (c) and (d) (presentation	n of extra claims)
		[]	37 C.F.R. Section 1.16(e) (surcharge for filing declaration on a date later than the filing date of the	ng the basic filing fee and/or ne application)
		[]	37 C.F.R. Section 1.17(a)(1)-(5) (extension fees po	ursuant to Section 1.136(a).
		[]	37 C.F.R. Section 1.17 (application processing fee	es)
		[]	37 C.F.R. Section 1.18 (issue fee at or before repursuant to 37 C.F.R. Section 1.311(b))	nailing of Notice of Allowance,

16.

	[X]	Credit Account No. 50-0850.	
	[]	Refund	
			SIGNATURE OF PRACTITIONER
Reg. N	o. 34,23	5	David S. Resnick (type or print name of practitioner)
Tel. No	o.: (617)	345-6057	101 Federal Street P.O. Address
Custon	ner No.:		Boston, MA 02110
[X]	(check applic division APPL CLAI	ation(s) (including an international onal or C-I-P application) and constant of the constant o	on in this transmittal claims the benefit of prior U.S. application entering the U.S. stage as a continuation, mplete and attach the ADDED PAGES FOR NEW BENEFIT OF PRIOR U.S. APPLICATION(S)
	[X]	Plus Added Pages for New Ap Application(s) Claimed	oplication Transmittal Where Benefit of Prior U.S. Number of pages added1
	[]	Plus Added Pages for Papers Refer	red to in Item 4 Above Number of pages added
	[]	Plus added pages deleting names of longer inventor(s) of the subject ma	f inventor(s) named on prior application(s) who is/are no atter claimed in this application. Number of pages added
	[]	Plus "Assignment Cover Letter Ace	companying New Application" Number of pages added

(New Application Transmittal--page 8 of 13)

Instructions as to Overpayment

[]	Statement Where No Further Pages Added						
	(if no further pages form a part of this Transmittal, then end this Transmittal with this page and check the following item)						
	[] This transmittal ends with this page.						

The House with the little of the world before the first the first

ADDED PAGE(S) FOR SPECIAL COMMENTS FOR APPLICATION TRANSMITTAL WHERE BENEFIT OF PRIOR U.S. APPLICATION(S)N IS CLAIMED

17.	Relate	back					
Α.	35 U.S.	35 U.S.C. 119(e)					
В.	35 U.S.	.C. 120, 121 and 365(c)					
	[X]	"This application is a					
		[X] continuation[] continuation-in-part[] divisional					
	of copending application(s)						
	[X]	application number 08/500,306 filed on July 10, 1995.					
U.S."	[]	International Application filed on and which designated					
filed or	[] n July 10	"The non-provisional application designed above, namely application number 08/500,306), 1995, claims benefit of U.S. Provisional Application(s) No(s).:					
APPL	ICATIO	ON NO(S).: FILING DATE					
	1	99					
	/						
	1	>					

Added page ____1___

Practitioner's Docket No. 953-45394-CPA-2C



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of: SCHLOM, et al.

Application No.:

Group No.:

Filed: Submitted herewith

Examiner:

For: GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC ANTIGEN (PSA)

Assistant Commissioner for Patents Washington, D.C. 20231

EXPRESS MAIL CERTIFICATE

"Express Mail" label number EL565095244US Date of Deposit 10/20/2000

I hereby state that the following attached papers or fees

New Application Transmittal

Copy of the Application No. 08/500,306 including 47 pages of Specification, Abstract, 2 pages of Sequence Listing, 2 sheets of Drawings;

Copy of Declaration and Power of Attorney;

Check in the sum of \$788.00; and

Return Receipt Postcard and Change of Attorney's Address in Application.

is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 C.F.R. section 1.10, on the date indicated above and is addressed to the Assistant Commissioner for Patents, Washington, D.C. 20231.

Patricia Turner

33421 Docket No.: 45394

GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC ANTIGEN (PSA)

FIELD OF THE INVENTION

The present invention relates generally to generation of cellular and humoral immune responses to a mammalian prostate-specific antigen (PSA).

BACKGROUND OF THE INVENTION

Cancer of the prostate is the most commonly diagnosed cancer in men and is the second most common cause of cancer death (Carter, et al., 1990; Armbruster, et al., 1993). If detected at an early stage, prostate cancer is potentially curable. However, a majority of cases are diagnosed at later stages when metastasis of the primary tumor has already occurred (Wang, et al., 1982). Even early diagnosis is problematic because not all individuals who test positive in these screens develop cancer. Present treatment for prostate cancer includes radical prostatectomy, radiation therapy, or hormonal therapy. No systemic therapy has clearly improved survival in cases of hormone refractory disease. With surgical intervention, complete eradication of the tumor is not always achieved and the observed re-occurrence of the cancer (12-68%) is dependent upon the initial clinical tumor stage (Zietman, et al., 1993). Thus, alternative methods of treatment including prophylaxis or prevention are desirable.

Prostate specific antigen (PSA) is a 240 amino acid member of the glandular kallikrein gene family. (Wang, et al., 1982; Wang, et al., 1979; Bilhartz, et al., 1991). PSA is a serine protease, produced by normal prostatic tissue, and secreted exclusively by the epithelial cells lining prostatic acini and ducts (Wang, et al., 1982; Wang, et al., 1979; Lilja, et al., 1993). Prostate specific antigen can be detected at low levels in the sera of healthy males without clinical evidence of prostate cancer.

25

30

5

10

However, during neoplastic states, circulating levels of this antigen increase dramatically, correlating with the clinical stage of the disease (Schellhammer, et al., 1993; Huang, et al., 1993; Kleer, et al., 1993; Oesterling, et al., 1991). Prostate specific antigen is now the most widely used marker for prostate cancer. The tissue specificity of this antigen makes PSA a potential target antigen for active specific immunotherapy (Armbruster, et al., 1993; Brawer, et al., 1989), especially in patients who have undergone a radical prostatectomy in which the only PSA expressing tissue in the body should be in metastatic deposits. Recent studies using in-vitro immunization have shown the generation of CD4 and CD8 cells specific for PSA (Peace et al., 1994; Correale et al., 1995). However, although weak natural killer cell responses have been occasionally documented in prostate cancer patients (Choe, et al., 1987), attempts to generate an in vivo immune response have met with limited success. For example, several attempts to actively immunize patients with prostate adenocarcinoma cells admixed with Bacillus Calmette-Guerin (BCG) have shown little or no therapeutic benefit (Donovan, et al., 1990). The ability to elicit an immune response as a result of exposure to PSA in vivo would be extremely useful.

Vaccinia virus has been used in the world-wide eradication of smallpox. This virus has been shown to express a wide range of inserted genes, including several tumor associated genes such as p97, HER-2/neu, p53 and ETA (Paoletti, et al., 1993). Other pox viruses that have been suggested as useful for expression of multiple genes include avipox such as fowl pox. Cytokines expressed by recombinant vaccinia virus include IL-1, IL-2, IL-5, IL-6, TNF- α and IFN- γ (Paoletti, et al., 1993). Recombinant pox viruses, for example vaccinia viruses, are being considered for use in therapy of cancer because it has been shown in animal models that the copresentation of a weak immunogen with the highly immunogenic poxvirus

10

proteins can elicit a strong immune response against the inserted gene product (Kaufman, et al., 1991; Paoletti, et al., 1993; Kantor, et al., 1992a; Kantor, et al., 1992b; Irvine, et al., 1993; Moss, et al., 1993). A recombinant vaccinia virus containing the human carcinoembryonic antigen gene has just completed phase 1 clinical trials in carcinoma patients with no evidence of toxicity other than that observed with the wild type smallpox vaccine (Kantor, et al., 1992b).

Currently, models for the evaluation of prostate therapeutics include the canine (McEntee, et al., 1987), and the Dunning rat (Isaacs, et al., 1986); neither of these models, however, are practical for the study of PSA-recombinant vaccines due to the very low homology of rat and canine PSA to human PSA (Karr, et al., 1995; Schroder, et al., 1982). In contrast, the prostate gland of the rhesus monkey is structurally and functionally similar to the human prostate (Wakui, et al., 1992). At the molecular level, there is 94% homology between either the amino acid or nucleic acid sequences of rhesus PSA (Gauthier, et al., 1993) and those sequences of human prostate specific antigen (Karr, et al., 1995; Lundwall, et al., 1987). Thus, human PSA is essentially an autoantigen in the rhesus monkey. Accordingly, the rhesus monkey can serve as a model for autologous anti-PSA immune reactions.

SUMMARY OF THE INVENTION

We have discovered that by using a recombinant viral vector, preferably a pox virus vector having at least one insertion site containing a DNA segment encoding prostate-specific antigen (PSA), or a cytotoxic T-cell eliciting epitope thereof, operably linked to a promoter capable of expression in the host, a specific humoral and cellular immune response to PSA can be generated. The method preferably comprises introducing a sufficient amount of the recombinant pox virus vector into a host to

30

10

PSA at periodic intervals thereafter. The additional PSA, or a cytotoxic T-cell eliciting epitope thereof, may be added by using a second pox virus vector from a different pox genus. In another embodiment, additional PSA can be added by contacting the host with PSA by a variety of other methods, including in one preferred embodiment adding PSA. The PSA may be formulated with an adjuvant or in a liposomal formulation.

In a further embodiment, an immune response to PSA can be generated by contacting the host initially with a sufficient amount of PSA, or a cytotoxic T-cell eliciting epitope thereof, to stimulate an immune response and at periodic intervals thereafter contacting the host with additional PSA. The additional PSA, or a cytotoxic T-cell generating fragment thereof, may be added using a pox virus vector as discussed above.

We have also discovered that human cytotoxic T-cells specific for PSA can be produced using a cytotoxic T-cell eliciting epitope of the PSA and that these cells have the ability to lyse PSA-expressing human prostate carcinoma cells.

As used herein the term "prostate specific antigen" includes the native protein whether purified from a native source or made by recombinant technology, as well as any polypeptide, mutein or portion derived therefrom that is capable of generating an immune response to a native conformationally correct PSA. For example, one can make conservative amino acid substitutions in the molecule without adversely affecting the ability to use the recombinant to generate an antibody that will also recognize native PSA.

30

5

10

The pox virus is preferably selected from the group of pox viruses consisting of suipox, avipox, capripox and orthopox virus. Preferred orthopox include vaccinia, rabbit pox and raccoon pox. Preferred avipox includes fowlpox, canary pox and pigeon pox. A more preferred avipox is fowlpox. The preferred suipox is swinepox.

Vaccinia viral vectors may elicit a strong antibody response. Thus while numerous boosts with vaccinia vectors are possible, its repeated use may not be preferred in certain instances. We have discovered that by using pox from different genera to boost, this sensitivity problem can be minimized. In accordance with the present invention, in order to avoid such problems, preferably, when the first or initial pox virus vector is vaccinia, the second and subsequent pox virus vectors are selected from the pox viruses from a different genus such as suipox, avipox, capripox or an orthopox immunogenically distinct from vaccinia.

Adjuvants include, for example, RIBI Detox, QS21, and incomplete Freund's adjuvant. Liposomal formulations can also be used.

Human cytotoxic T-cells specific for PSA produced in accordance with the present invention can be isolated from a human host. These cells can be used in drug assays, used to map cytotoxic T-cells eliciting antigen epitopes or in adoptive cell therapy.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows a Western blot of PSA from rV-PSA infected BSC-40 cells. Lanes 2-4 are extracts from supernatant fluid from cells infected overnight with rV-PSA at an MOI of 1, while Lanes 7-9 are extracts from the corresponding infected cells. Lanes 1 and 7 are supernatant extracts and cell extracts from V-Wyeth infected cells. Blot was developed using a

10

specific MAb for human PSA. This blot illustrates that cells infected with rV-PSA authentically express and secrete the 33 kD PSA protein.

Figures 2A, 2B and 2C show the manifestation of rV-PSA immunization. In Figure 2A, the area of lesions was measured 7 days following each inoculation of rhesus monkeys with either V-Wyeth (open circles) or rV-PSA (closed circles). In Figure 2B, the duration of the lesion was monitored as time of scab disappearance. In Figure 2C, the extent of lymph node swelling was recorded and characterized as very swollen (3+), i.e., more than two axillary nodes swollen; swollen (2+), i.e., one or two nodes easily palpable; marginally swollen (1+), i.e., one node was barely palpable; or not swollen (0), 7 days following inoculation with vaccinia virus. Each symbol represents one monkey.

DETAILED DESCRIPTION OF THE INVENTION

We have induced an immune response specific to PSA in the rhesus monkey model by placing the PSA gene into a recombinant viral vector, i.e, a pox vector such as vaccinia virus.

Additionally, an immune response to PSA can be generated by contacting the host initially with a sufficient amount of PSA, or a cytotoxic T-cell eliciting epitope thereof, to stimulate an immune response and at periodic intervals thereafter contacting the host with additional PSA. The additional PSA, or a cytotoxic T-cell generating fragment thereof, may be added using a pox virus vector.

A DNA fragment encoding the open reading frame of human PSA can be obtained, for example, from total RNA extracted from the human metastatic prostate adenocarcinoma cell line, LNCaP.FGC (CRL 1740, American Type Cell Culture (ATCC), Rockville, MD) by reverse

30

10

transcriptase PCR using PSA specific oligonucleotide primers 5'
TCTAGAAGCCCCAAGCTTACCACCTGCA 3' (SEQ. ID. NO.:1), 5'
TCTAGATCAGGGGTTGGCCACGATGGTCCTTGATCCACT 3' (SEQ. ID. NO.:2). The nucleotide sequence of the PSA cDNA has been published (Lundwall, et al., 1987).

Recombinant human PSA can be obtaining using a baculovirus expression system in accordance with the method of Bei et al., *J. Clin. Lab. Anal.*, 9:261-268 (1995), the disclosure of which is herein incorporated by reference.

Viral vector

Basic techniques for preparing recombinant DNA viruses containing a heterologous DNA sequence encoding the carcinoma self-associated antigen or cytotoxic T-cell eliciting epitope are known to the skilled artisan and involve, for example, homologous recombination between the viral DNA sequences flanking the DNA sequence in a donor plasmid and homologous sequences present in the parental virus (Mackett, et al., *Proc. Natl. Acad. Sci. USA* 79:7415-7419 (1982)). For example, recombinant viral vectors such as a pox viral vector can be used in delivering the gene. The vector can be constructed for example by steps known in the art, e.g. analogous to the methods for creating synthetic recombinants of the fowlpox virus described in U.S. Patent No. 5,093,258, the disclosure of which is incorporated herein by reference. Other techniques include using a unique restriction endonuclease site that is naturally present or artificially inserted in the parental viral vector to insert the heterologous DNA.

Pox viruses useful in practicing the present invention include orthopox, suipox, avipox and capripox virus.

30

10

Orthopox include vaccinia, ectromelia and raccoon pox. The preferred orthopox is vaccinia.

Avipox includes fowlpox, canary pox and pigeon pox. The preferred avipox is fowlpox.

Capripox include goatpox and sheeppox.

A preferred suipox is swinepox.

Other viral vectors that can be used include other DNA viruses such as herpes virus and adenoviruses, and RNA viruses such as retroviruses and polio.

For example, the DNA gene sequence to be inserted into the virus can be placed into a donor plasmid, e.g., an *E. coli* plasmid construct, into which DNA homologous to a section of DNA such as that of the insertion site of the poxvirus where the DNA is to be inserted has been inserted. Separately the DNA gene sequence to be inserted is ligated to a promoter. The promoter-gene linkage is positioned in the plasmid construct so that the promoter-gene linkage is flanked on both ends by DNA homologous to a DNA sequence flanking a region of pox DNA which is the desired insertion region. With a parental pox viral vector, a pox promoter is used. The resulting plasmid construct is then amplified by growth within *E. coli* bacteria and isolated. Preferably, the plasmid also contains an origin of replication such as the *E. coli* origin of replication, and a marker such as an antibiotic resistance gene for selection and propagation in *E. coli*.

Second, the isolated plasmid containing the DNA gene sequence to be inserted is transfected into a cell culture, e.g., chick embryo fibroblasts,

30

30

5

10

along with the parental virus, e.g., poxvirus. Recombination between homologous pox DNA in the plasmid and the viral genome respectively results in a recombinant poxvirus modified by the presence of the promoter-gene construct in its genome, at a site which does not affect virus viability.

As noted above, the gene is inserted into a region (insertion region), in the virus which does not affect virus viability of the resultant recombinant virus. The skilled artisan can readily identify such regions in a virus by, for example, randomly testing segments of virus DNA for regions that allow recombinant formation without seriously affecting virus viability of the recombinant. One region that can readily be used and is present in many viruses is the thymidine kinase (TK) gene. For example, the TK gene has been found in all pox virus genomes examined [leporipoxvirus: Upton, et al., J. Virology, 60:920 (1986) (shope fibroma virus); capripoxvirus: Gershon, et al., J. Gen. Virol., 70:525 (1989) (Kenya sheep-1); orthopoxvirus: Weir, et al., J. Virol., 46:530 (1983) (vaccinia); Esposito, et al., Virology, 135:561 (1984) (monkeypox and variola virus); Hruby, et al., PNAS, 80:3411 (1983) (vaccinia); Kilpatrick, et al., Virology, 143:399 (1985) (Yaba monkey tumor virus); avipoxvirus: Binns, et al., J. Gen. Virol. 69:1275 (1988) (fowlpox); Boyle, et al., Virology, 156:355 (1987) (fowlpox); Schnitzlein, et al., J. Virological Methods, 20:341 (1988) (fowlpox, quailpox); entomopox (Lytvyn, et al., J. Gen. Virol. 73:3235-3240 (1992)].

In vaccinia, in addition to the TK region, other insertion regions include, for example, the HindIII M fragment.

In fowlpox, in addition to the TK region, other insertion regions include, for example, the BamHI J fragment [Jenkins, et al., AIDS Research

5

20 m

25

30

and Human Retroviruses 7:991-998 (1991)] the EcoRI-HindIII fragment, EcoRV-HindIII fragment, BamHI fragment and the HindIII fragment set forth in EPO Application No. 0 308 220 A1. [Calvert, et al., J. of Virol. 67:3069-3076 (1993); Taylor, et al., Vaccine 6:497-503 (1988); Spehner, et al., (1990) and Boursnell, et al., J. of Gen. Virol. 71:621-628 (1990)].

In swinepox preferred insertion sites include the thymidine kinase gene region.

In addition to the requirement that the gene be inserted into an insertion region, successful expression of the inserted gene by the modified poxvirus requires the presence of a promoter operably linked to the desired gene, i.e., in the proper relationship to the inserted gene. The promoter must be placed so that it is located upstream from the gene to be expressed. Promoters are well known in the art and can readily be selected depending on the host and the cell type you wish to target. For example in poxviruses, pox viral promoters should be used, such as the vaccinia 7.5K, 40K or fowlpox promoters such as FPV C1A. Enhancer elements can also be used in combination to increase the level of expression. Furthermore, the use of inducible promoters, which are also well known in the art, in some embodiments are preferred.

A specific immune response for PSA can be generated by administering between about 105-109 pfu of the recombinant pox virus, constructed as discussed above to a host, more preferably one uses 107 pfu. The preferred host is a human. At least one interval thereafter, which is preferably one to three months later, the immune response is boosted by administering additional antigen to the host. More preferably there is at least a second "boost" preferably one to three months after the first boost. The antigen may be administered using the same pox virus vector. The

10

30

25

antigen may preferably be administered using a second pox virus vector from a different pox genera, or may be administered directly using, for example, an adjuvant or liposome. Cytokines, e.g., IL-2, IL-6, IL-12 or costimulatory molecules, e.g., B7.1, B7.2, may be used as biologic adjuvants and can be administered systemically to the host or co-administered via insertion of the genes encoding the molecules into the recombinant pox vector.

Adjuvants include, for example, RIBI Detox (Ribi Immunochemical), QS21 and incomplete Freund's adjuvant.

Generation of Cytotoxic T-Cells

Cytotoxic T-cells specific for PSA can be established from peripheral blood mononuclear cells (PBMC) obtained from a host immunized as discussed above. For example, PBMC can be separated by using Lymphocyte Separation Medium gradient (Organon Teknika, Durham, NC, USA) as previously described [Boyum, et al., Scand J. Clin Lab Invest 21: 77-80 (1968)]. Washed PBMC are resuspended in a complete medium, for example, RPMI 1640 (GIBCO) supplemented with 10% pool human AB serum (Pel-Freeze Clinical System, Brown Dear, WI, USA), 2mM glutamine, 100 U/ml penicillin and 100 μ g/ml of streptomycin (GIBCO). PBMC at a concentration of about 2 x 105 cells in complete medium in a volume of, for example, 100 μ l are added into each well of a 96-well flat-bottom assay plate (Costar, Cambridge, MA, USA). The antigen or peptides are added into the cultures in a final concentration of about 50 $\mu \mathrm{g/ml}$ and incubated at 37°c in a humidified atmosphere containing 5% CO2 for 5 days. After removal of peptide containing media, the cultures are provided with fresh human IL-2 (10U/ml) after 5 days and replenished with IL-2 containing medium every 3 days. Primary cultures are restimulated with the same peptide (50 μ g/ml) on day 16. 5 x 10⁵ irradiated (4,000 rad)

10

autologous PBMC are added in a volume of about 50 μ l complete medium as antigen-presenting cells (APC). About five days later, the cultures are provided with human IL-2 containing medium as described previously. Cells are restimulated for 5 days at intervals of 16 days.

Epitope mapping

The cytotoxic T-cells of the present invention can be used to determine the epitope of the PSA that elicits a cytoxic T-cell. For example, one can cut the PSA into numerous peptide fragments. Alternatively, the fragments can be chemically synthesized. Cytotoxic T-cells can then be plated and different fragments added to different wells. Only T-cells which recognize one of the pre-selected peptide fragments as an epitope will continue to expand, thereby permitting ready identification.

These fragments can then be used to elicit cytotoxic T-cell instead of using the whole protein. Additionally, one can prepare other fragments containing the epitope to enhance its ability to elicit a cytoxic T-cell response. Modifications to these fragments are well known in the art and include the use of conjugates, specific amino acid residues such as cystines, etc.

Drug Assay

The cytotoxic T-cell can also be used to screen for compounds which enhance the ability of the antigen to create a cytotoxic T-cell response. For example, cytotoxic T-cells can be incubated with a selected epitope, for example, in a microtiter plate. The compound to be tested, e.g. a drug, is then added to the well and the growth of the T-cells is measured. T-cell expansion indicates that the test compound enhances the T-cell response. Such compounds can be further evaluated.

10

Therapy

The cytotoxic T-cell can be cultured to amplify its number and then injected back into the host by a variety of means. Generally, between 1 x 10⁵ and 2 x 10¹¹ cytotoxic T-cells per infusion are administered in, for example, one to three infusions of 200 to 250 ml each over a period of 30 to 60 minutes. After the completion of the infusions, the patient may be treated with recombinant interleukin-2 with a dose of 720,000 IU per kilogram of body weight intravenously every eight hours; some doses can be omitted depending on the patient's tolerance for the drug. In addition, after infusion, additional antigen or fragments containing T-cell eliciting epitope(s) may be administered to the patient to further expand the T-cell number. The antigen or epitope may be formulated with an adjuvant and/or may be in a liposomal formulation.

The cytotoxic T-cells can also be modified by introduction of a viral vector containing a DNA encoding TNF and reintroduced into a host in an effort to enhance the anti-tumor activity of the cells. Other cytokines can also be used.

The recombinant vector can be administered using any acceptable route, including, for example, scarification and injection, e.g., intradermal, subcutaneous, intramuscular, intravenous or intraperitoneal.

For parenteral administration, the recombinant vectors will typically be injected in a sterile aqueous or non-aqueous solution, suspension or emulsion in association with a pharmaceutically-acceptable carrier such as physiological saline.

REFERENCE EXAMPLE 1 CONSTRUCTION OF VECTORS

30

5

10

Pox Viruses

A number of pox viruses have been developed as live viral vectors for the expression of heterologous proteins (Cepko et al., *Cell* 37:1053-1062 (1984); Morin et al., *Proc. Natl. Acad. Sci. USA* 84:4626-4630 (1987); Lowe et al., *Proc. Natl. Acad. Sci. USA*, 84:3896-3900 (1987); Panicali & Paoletti, *Proc. Natl. Acad. Sci. USA*, 79:4927-4931 (1982); Mackett et al., *Proc. Natl. Acad. Sci. USA*, 79:7415-7419 (1982)). Representative fowlpox and swinepox virus are available through the ATCC under accession numbers VR-229 and VR-363, respectively.

DNA Vectors For In Vivo Recombination With A Parent Virus

Genes that code for desired carcinoma associated antigens are inserted into the genome of a pox virus in such a manner as to allow them to be expressed by that virus along with the expression of the normal complement of parent virus proteins. This can be accomplished by first constructing a DNA donor vector for *in vivo* recombination with a pox virus.

In general, the DNA donor vector contains the following elements:

- (i) a prokaryotic origin of replication, so that the vector may be amplified in a prokaryotic host;
- (ii) a gene encoding a marker which allows selection of prokaryotic host cells that contain the vector (e.g., a gene encoding antibiotic resistance);
- (iii) at least one gene encoding a desired protein located adjacent to a transcriptional promoter capable of directing the expression of the gene; and

(iv) DNA sequences homologous to the region of the parent virus genome where the foreign gene(s) will be inserted, flanking the construct of element (iii).

Methods for constructing donor plasmids for the introduction of

5

10

multiple foreign genes into pox virus are described in W091/19803, the techniques of which are incorporated herein by reference. In general, all DNA fragments for construction of the donor vector, including fragments containing transcriptional promoters and fragments containing sequences homologous to the region of the parent virus genome into which foreign genes are to be inserted, can be obtained from genomic DNA or cloned DNA fragments. The donor plasmids can be mono-, di-, or multivalent (i.e., can contain one or more inserted foreign gene sequences).

The donor vector preferably contains an additional gene which encodes a marker which will allow identification of recombinant viruses containing inserted foreign DNA. Several types of marker genes can be used to permit the identification and isolation of recombinant viruses. These include genes that encode antibiotic or chemical resistance (e.g., see Spyropoulos et al., *J. Virol.*, 62:1046 (1988); Falkner and Moss., *J. Virol.*, 62:1849 (1988); Franke et al., *Mol. Cell. Biol.*, 5:1918 (1985), as well as genes such as the *E. coli lac*Z gene, that permit identification of recombinant viral plaques by colorimetric assay (Panicali et al., *Gene*, 47:193-199 (1986)).

25

30

Integration Of Foreign DNA Sequences Into The Viral Genome And Isolation Of Recombinants

Homologous recombination between donor plasmid DNA and viral DNA in an infected cell results in the formation of recombinant viruses that incorporate the desired elements. Appropriate host cells for *in vivo* recombination are generally eukaryotic cells that can be infected by the

10

25

30

virus and transfected by the plasmid vector. Examples of such cells suitable for use with a pox virus are chick embryo fibroblasts, HuTK143 (human) cells, and CV-1 and BSC-40 (both monkey kidney) cells. Infection of cells with pox virus and transfection of these cells with plasmid vectors is accomplished by techniques standard in the art (Panicali and Paoletti, U.S. Patent No. 4,603,112, W089/03429).

Following *in vivo* recombination, recombinant viral progeny can be identified by one of several techniques. For example, if the DNA donor vector is designed to insert foreign genes into the parent virus thymidine kinase (TK) gene, viruses containing integrated DNA will be TK and can be selected on this basis (Mackett et al., *Proc. Natl. Acad. Sci. USA*, 79:7415 (1982)). Alternatively, co-integration of a gene encoding a marker or indicator gene with the foreign gene(s) of interest, as described above, can be used to identify recombinant progeny. One preferred indicator gene is the *E. coli lac*Z gene: recombinant viruses expressing β -galactosidase can be selected using a chromogenic substrate for the enzyme (Panicali et al., *Gene*, 47:193 (1986)).

Characterizing The Viral Antigens Expressed By Recombinant Viruses

Once a recombinant virus has been identified, a variety of methods can be used to assay the expression of the polypeptide encoded by the inserted gene. These methods include black plaque assay (an *in situ* enzyme immunoassay performed on viral plaques), Western blot analysis, radioimmunoprecipitation (RIPA), and enzyme immunoassay (EIA).

EXAMPLE I

Generation of PSA Specific Immune Response

Materials and Methods

15 In the state of the state of

5

10

Recombinant Vaccinia Virus

A 786 bp DNA fragment encoding the entire open reading frame of human prostate specific antigen was amplified by reverse transcriptase PCR (GeneAmp RNA PCR Kit, Perkin Elmer, Norwalk, CT) from total RNA extracted from the human metastatic prostate adenocarcinoma cell line, LNCaP.FGC (CRL 1740, American Type Culture Collection (ATCC), Rockville, MD). The predicted amino acid sequence derived from the PSA coding sequence was shown to be nearly identical to the published sequence (Lundwall, et al., 1987), differing only in a change from asparagine to tyrosine at position 220. The PSA DNA fragment, containing the entire coding sequence for PSA, 41 nucleotides of the 5' untranslated region, and 520 nucleotides of the 3' untranslated region, was inserted into the Xba I restriction endonuclease cleavage site of the vaccinia virus transfer vector pT116. The resulting plasmid, designated pT1001, contains the PSA gene under the control of the vaccinia virus 40K promoter (Gritz, et al. 1990) and the E. coli lacZ gene under the control of the fowlpox virus C1 promoter (Jenkins, et al., 1991). The foreign genes are flanked by DNA sequences from the Hind III M region of the vaccinia genome. A plaque-purified isolate from the Wyeth (New York City Board of Health) strain of vaccinia was used as the parental virus in the construction of the recombinant vaccinia virus. The generation of recombinant vaccinia virus was accomplished via homologous recombination between vaccinia sequences in the Wyeth vaccinia genome and the corresponding sequences in pT1001 in vaccinia-infected RK₁₃ cells (CCL 37, ATCC) transfected with pT1001. Recombinant virus was identified using a chromogenic assay, performed on viral plaques in situ, that detects expression of the lacZ gene product in the presence of halogenated indolyl-beta-D-galactoside (Bluo-gal), as described previously (Panacali, et al., 1986). Appropriate blue recombinant viruses were purified by four rounds of plaque-purification. Virus stocks were prepared

30

10

by clarifying infected RK₁₃ cell lysates followed by centrifugation through a 36% sucrose cushion.

Characterization of Recombinant Virus

Southern Analysis of DNA recombination

The recombinant vaccinia genome was analyzed by viral DNA extraction, restriction endonuclease digestion with Hind III, and Southern blotting as previously described (Kaufman et al., 1991).

Western Analysis of protein expression

Confluent BSC-40 cells were infected with either parental wild type vaccinia virus (designated V-Wyeth) or recombinant vaccinia-PSA (designated rV-PSA) at an MOI of 1 in Dulbecco's Modified Eagle's Medium containing 2% fetal bovine serum. After an overnight infection, the medium was removed from the cells, and an aliquot was methanol precipitated to assay for the presence of secreted PSA. The infected cells were lysed in hypotonic lysis buffer (150 mM NaCI, 0.05% EDTA, 10 mM KCI, 1 mM PMSF) and then sonicated. Cell lysates and culture media were electrophoresed on an SDS-10% acrylamide gel. The proteins were transblotted to nitrocellulose, and the blot was incubated with a rabbit antibody specific for PSA (P0798, Sigma Chemical Co., St. Louis, MO) for 4 hours at ambient temperature, washed, and then incubated with goat anti-rabbit phosphatase-labeled secondary antibody (AP, Kirkegaard & Perry Laboratories, Gaithersburg, MD) and developed according to the manufacture's instructions.

Generation of B-cell lines

Monkey autologous B lymphoblastoid cell lines (BLCL) were established by infecting 1x10⁵ freshly isolated PBMCs in 100 ml of RPMI 1640 supplemented with L-glutamine, gentamicin, and 10% FCS

30

10

(Biofluids, Rockville, MD) with 100 ml supernatant from S594 cells (kindly provided by Dr. M. D. Miller, Harvard Medical School, New England Regional Primate Research Center, Southborough, MA), which contains the baboon herpesvirus *Herpes papio*, in a 96 well, flat-bottomed plate (Costar, Cambridge, MA). Following transformation, cells were expanded, and media changed once weekly.

Immunization of Monkeys

Twelve juvenile male rhesus monkeys (*Macaca mulatta*), ages 1 to 2 years, were assigned to three vaccination groups of four animals each. One animal from each group was prostatectomized. Animals were immunized 3 times on days 1, 29, and 57. Doses of either 1 x 10⁷ or 1 x 10⁸ PFU of rV-PSA were administered to 4 animals by skin scarification. V-Wyeth (1 x 10⁸ PFU) was administered to 4 animals as controls. The animals were housed and maintained at the Toxicology Research Laboratory, University of Illinois at Chicago (TRL/UIC) in accordance with the guidelines of the National Cancer Institute Animal Care and Use Committee and the Guide for the Care and Use of Laboratory Animals (Department of Health and Human Services Publication NIH 85-23, revised 1985 by the FDA Center for Biologics Evaluation and Research Office of Biological Product Review, Division of Product Quality Control, Pathology and Primatology Laboratory, Bethesda, MD).

Toxicology

25

Physical examinations were performed on ketamine (Ketamine® HCl, 10 mg/kg l.M.) sedated animals. Rectal temperatures and weights were recorded for each monkey on a weekly basis. The vaccination site was observed and erythema and swelling were measured by caliper. Each animal was examined for regional lymphadenopathy, hepatomegaly, and splenomegaly. Any other gross abnormalities were also recorded.

10

Blood was obtained by venipuncture from the femoral vein of ketamine sedated animals before and after each immunization. A complete blood count, differential, hepatic and renal chemistry evaluation was performed on each monkey by TRL/UIC. Results were compared to normal primate values (Kantor *et al.*, 1992b). Circulating levels of PSA before and after immunization were analyzed by radioimmunoassay (TandemTM, Hybritech, San Diego, CA).

Measurement of Antibody Titers

Prior to each immunization and 2 weeks following each immunization, anti-PSA antibody was quantified by ELISA. Microtiter plates were coated with purified PSA (100 ng/well, Calbiochem, La Jolla, CA), ovalbumin (100 ng/well, Sigma), or 1x10⁷ PFU/well UV-inactivated V-Wyeth in PBS. The plates were blocked with 2% BSA in PBS, dried, and stored at -20° C until used. The plates were incubated with serum diluted 1:5, as well as a monoclonal antibody for PSA (DAKO M750, Denmark) as a standard control, for 24 hours at 4° C. Plates were washed several times with PBS containing 1% BSA, and incubated at 37° C for 45 min with horseradish peroxidase-conjugated goat anti-human IgG or IgM heavy chain specific antiserum (1:8000) (Southern Biotechnology Associates, Birmingham, AL) and antibody detected by HRP substrate system (Kirkegaard & Perry Laboratories, Gaithersburg, MD) according to the manufacture's instructions. The absorbance of each well was read at 405 nm using a Bio-Tek EL310 microplate ELISA reader (Winooski, VT).

Lymphoproliferative Assay

Autologous monkey BLCL were plated at a density of 3 x 10⁶ cells/well in 24 well plates with 160 mg/well purified PSA (Fitzgerald, Concord, MA) or 160 mg/well ovalbumin (Sigma) at 37° C for 24 hours. Cells were then *y*-irradiated (14000 rad), harvested, washed and

30

10

suspended at a final concentration of 1 x 10⁷/ml. Fresh monkey PBMCs from heparinized blood, 6 weeks to 7 months after the last immunization, were isolated on lymphocyte separation medium (Organon Teknika, West Chester, PA). Lymphoproliferative responses were evaluated by coculturing 1.5 x 10⁵ cells with 5 x 10⁵ cells/well of autologous BLCL in 0.2 ml of RPMI 1640 supplemented with 10% heat-inactivated fetal calf serum in flat-bottomed 96 well plates (Costar) for 5 days. PBMCs were cultured with 2 x 10⁷ PFU/ml UV-inactivated V-Wyeth as a recall antigen or 2 mg/ml Con-A as positive controls. Cells were labeled for the final 12-18 h of the incubation with 1 mCi/well [³H]thymidine (New England Nuclear, Wilmington, DE) and harvested with a PHD cell harvester (Cambridge Technology, Cambridge, MA). The incorporated radioactivity was measured by liquid scintillation counting (LS 6000IC; Beckman, Duarte, CA). The results from triplicate wells were averaged and are reported as mean ± SEM.

Results

Generation and Characterization of Recombinant Virus

The cDNA fragment encoding the open reading frame of human PSA was obtained by reverse transcriptase PCR using PSA specific oligonucleotide primers 5' TCTAGAAGCCCCAAGCTTACCACCTGCA 3' (SEQ. ID. NO.:1), 5'

TCTAGATCAGGGGTTGGCCACGATGGTGTCCTTGATCCACT 3' (SEQ. ID. NO.:2), and ligated into the vaccinia virus transfer vector pT106. This vector contains a strong vaccinia virus early/late promoter (designated P40) upstream of the multiple cloning site to drive the synthesis of the inserted gene product. The ligation and orientation of the PSA DNA fragment, as well as promoter position were verified by PCR and sequencing. The chimeric vector construct was inserted into the vaccinia virus genome Hind III M site by homologous recombination as previously

30

5

reported (Kaufman, et al., (1991)), and confirmed by Southern analysis probing with ³²P radiolabeled DNA corresponding to PSA sequences and vaccinia sequences in the Hind III M region (data not shown). The entire cDNA sequence of PSA in the vaccinia virus clone was shown to be nearly identical to the published sequences (Lundwall, et al., 1987).

Expression of recombinant protein was confirmed by western blot analysis of supernatant fluids and protein extracts from rV-PSA infected BSC-40 cells. These cells are routinely used for the evaluation of recombinant vaccinia products (Moss, et al., 1993). Incubation of cell supernatant blots from rV-PSA infected cells with rabbit anti-PSA antibody revealed a single immunoreactive polypeptide of approximately 33,000 daltons (Figure 1, lanes 2-4). Similarly, incubation of protein extract blots from rV-PSA infected cells revealed a single band of the same molecular weight (Figure 1, lanes 7-9). This is consistent with the predicted size of the PSA molecule (Armbruster, et al., 1993; Wang, et al., 1982). Cell supernatant blots (lane 1) or protein extract blots (lane 6) from cells infected with parental strain V-Wyeth remained negative for expression of PSA. These results thus demonstrate that a recombinant vaccinia virus can faithfully express the human PSA gene product.

Rhesus Monkey Model

The prostate gland of the rhesus monkey is structurally and functionally similar to the human prostate (Wakui, et al., 1992). At the molecular level, there is 94% homology between both the amino acid and nucleic acid sequences of rhesus PSA (Gauthier, et al., 1993) and human prostate specific antigen (Karr, et al., 1995; Lundwall, et al., 1987). Human PSA is essentially an autoantigen in the rhesus monkey.

Experimental Design

10

25

30

Table 1 delineates the protocol used in the immunization of 12 rhesus monkeys with either rV-PSA or the control V-Wyeth by skin scarification. Three groups of 4 animals were immunized with either rV-PSA at 1x10⁷ PFU/dose, rV-PSA at 1x10⁸PFU/dose, or V-Wyeth at 10⁸PFU/dose 3 times at 4 week intervals. These doses were chosen to ascertain the maximum tolerated dose for safety as well as to obtain maximum humoral and cell-mediated responses to PSA.

The rhesus monkeys were divided into 3 groups: high dose V-Wyeth, low dose rV-PSA, and high dose rV-PSA. One animal in each group was surgically prostatectomized to parallel two situations with regard to potential therapy in humans: (a) prostate intact, with primary and/or metastatic disease; or (b) patients prostatectomized with prostate cancer metastatic deposits. The presence of an intact prostate gland could conceivably serve as an antigen 'sink', either inducing anergy through persistence of antigen, or masking immunological effects by sequestering reactive cells or antibodies.

Physical Consequence of Immunization

The area of the lesions induced by rV-PSA or V-Wyeth was analyzed 7 days following each inoculation. In general, more induration was seen after the first inoculation, compared to the second inoculation (Figure 2A). After the third inoculation, there was no swelling of the vaccination site. The duration of the lesion following each immunization was shorter after each inoculation (Figure 2B). Regional lymph node swelling following vaccination was greater in most monkeys following the first immunization, compared to the second, or third immunization (Figure 2C). In general, no differences were seen in these parameters with the use of rV-PSA or V-Wyeth. Monkeys receiving V-Wyeth were compared with those receiving rV-PSA with respect to constitutional symptoms. Mild temperature

30

5

10

elevations were seen in all animals following vaccination. There was no evidence of weight loss, hepatomegaly or splenomegaly in any of the animals, and there was no differences between V-Wyeth or rV-PSA treated animals (data not shown). Animals were tested for complete blood count, differential, and hepatic and renal chemistries. Complete blood counts remained within normal limits throughout the study in both V-Wyeth and rV-PSA immunized animals (Table 2). Hepatic and renal functions were assessed prior to immunization and 12 weeks following primary immunization (Table 3). Parameters analyzed included alkaline phosphatase, blood urea nitrogen, alanine aminotransferase, aspartate aminotransferase, lactate dehydrogenase, and creatine and creatine kinase levels. There was no significant difference between animals receiving V-Wyeth or rV-PSA. There was no detectable PSA in the circulation of any of these monkeys after any immunization (detection limit was 0.1 ng/ml). At this time, which is 54 weeks post all immunizations, no toxicities were observed in monkeys of any of the groups, including those which were prostatectomized.

PSA Specific Humoral Responses

As indicated in Table 1, monkeys 1-4 were administered V-Wyeth while monkeys 5-12 were administered rV-PSA. Sera from each of these monkeys were analyzed by ELISA for immunoreactivity to PSA or UV-inactivated V-Wyeth, and ovalbumin as control antigen. Sera obtained from monkeys prior to vaccination were negative for reactivity to PSA (Table 4, PI). Fifteen days following primary immunization, monkeys in both the 1x10⁸ and 1x10⁷ dose rV-PSA groups developed low titer IgM antibodies specific for PSA (titers were determined at a 1:5 serum dilution). Although other isotopes of antibody were analyzed (IgG, IgA, IgM), only IgM was induced by rV-PSA throughout the observation period of 270 days. The antibody titers decreased over the 4 weeks prior to the

10

next inoculation. Prior to the second vaccination on day 29, 3 of 4 animals in the 1x10⁷rV-PSA group remained positive for PSA antibody, while 4 of 4 animals remained positive in the 1x10⁸rV-PSA group. Anti-PSA antibody titers increased after the second vaccination on day 29, but remained static after the third vaccination on day 57. By 270 days after the primary immunization, all animals were negative for PSA IgM antibody. Monkeys remained negative for IgG specific for PSA throughout the observation period (data nor shown). There was no correlation between rV-PSA dose and anti-PSA IgM titer, nor was there any apparent effect of prostatectomy. All monkey sera were negative for IgG or IgM to ovalbumin at all time points; as a positive control, however, the IgG titer in all three treatment groups to vaccinia virus was greater than 1:2000 as early as 29 days after the primary immunization (data not shown).

In general, vaccinia virus is a weak human pathogen (Paoletti *et al.*, 1993). Following vaccination, local erythema, induration, low-grade fever, and regional lymphadenopathy are common. The virus replicates in the epidermal cells of the skin and the virus is usually cleared within 14 days. All monkeys, whether given V-Wyeth or rV-PSA, exhibited the usual low grade constitutional symptoms of a vaccinia virus infection (Figure 2). There was no evidence of any adverse effects as indicated by changes in blood counts, differentials, hepatic and renal chemistries (Tables 2-3). The monkeys appeared healthy, without any physical signs of toxicity, throughout the 54 weeks of observation.

Although the rV-PSA construct was unable to elicit an anti-PSA IgG response, PSA specific IgM responses were noted in all rV-PSA immunized monkeys regardless of dose level (Table 4). These antibody responses were of low titer, short lived and could not be boosted, indicating induction of a primary response but not memory B-cells or affinity maturation.

PSA Specific Lymphoproliferative Assay

PSA specific T-cell responses in monkeys immunized with rV-PSA or V-Wyeth were analyzed using a lymphoproliferative assay. As seen in Table 5, the PBMCs from all monkeys analyzed responded, regardless of whether they received rV-PSA or V-Wyeth, to the lymphocyte mitogen concanavalin-A, as well as with the recall antigen UV-inactivated V-Wyeth. Differential responses to PSA *versus* medium alone or ovalbumin were seen in 1 animal (number 6) in the 1 x 10⁷ PFU rV-PSA group. All PBMCs from animals in the 1 x 10⁸ PFU rV-PSA group, however, responded to PSA in this assay. This experiment was repeated 5 times with similar results and data shown in Table 5 is from PBMCs isolated from monkeys 270 days after the primary immunization. No differences in PSA specific T-cell responses were seen in the prostatectomized monkeys.

To investigate cell mediated responses to the administration of rV-PSA, lymphoproliferative assays were performed using PBMCs from animals receiving the recombinant vaccine. One of four monkeys receiving the lower dose of rV-PSA (1 x 10⁷ PFU) and four of four receiving the higher dose (1 x 10⁸ PFU) maintained specific T-cell responses to PSA protein up to 270 days following primary immunization as indicated by the lymphoproliferative assay (Table 5). Prostatectomy appeared to have no effect on either the humoral or cellular responses of monkeys receiving rV-PSA. Evidence of PSA specific T-cell responses in monkeys lacking mature antibody isotopes could be due to two distinct events following vaccination with rV-PSA: a T-cell independent event, leading to IgM production, and a T-cell dependent event, leading to specific lymphoproliferative responses.

Table 1 Inoculation protocol of rhesus monkeys with the PSA recombinant and wild-type vaccinia virus

20.2		J L	
Monkey	Prostate	Immunogen	Dose* (PFU)
1	Yes	V-Wyeth	1x108
2	Yes	V-Wyeth	1x108
3	Yes	V-Wyeth	1x10 ⁸
4	No	V-Wyeth	1x10 ⁸
5	Yes	rV-PSA	1x10 ⁷
6	Yes	rV-PSA	$1x10^{7}$
7	Yes	rV-PSA	$1x10^{7}$
8	No	rV-PSA	1×10^7
9	Yes	rV-PSA	1x108
10	Yes ·	rV-PSA	1x10 ⁸
11	Yes	rV-PSA	1x10 ⁸
12	No	rV-PSA	1x10 ⁸

^{*} All animals received 3 immunizations at 4 week intervals.

Table 2 Mean WBC count, hematocrit, and differential count in rhesus monkeys receiving recombinant or wild-type vaccine

		V-Wyeth (n=4)	th (n=4)	rV-PSA (n=8)	(n=8)
Tect	Normal ranges	Before immunization ^a	Before After immunization ^a immunization ^b	Before immunization	After immunization
	7-15 x 10 ³	5.0 ± 0.8	5.1 ± 0.5	5.2 ± 0.7	5.8 ± 0.9
WDC	33-43	37.4 ± 0.2	37.0 ± 0.1	37.8 ± 0.4	37.0 ± 0.5
riematociii (voi. 70)	17 × 103	2.8 ± 0.7	3.9 ± 0.5	2.2 ± 0.4	3.5 ± 0.8
Lymphocytes	01 V /-I	2.0 ± 0.2	0.78 ± 0.2	2.9 ± 0.6	1.9 ± 0.3
SEGS ^c (%)) ×-	0.1 ± 0.05	0.2 ± 0.04	0.1 ± 0.04	0.2 ± 0.50
Monocytes (%)	8-O	0.1 ± 0.02	0.2 ± 0.10	0.1 ± 0.03	0.1 ± 0.02
Cosmobines (10)					

immunization a 1 week prior to primary immunization b 12 weeks following primary immuniza

c Segmented lymphocytes

Table 3 Mean serum chemistry values in rhesus monkeys receiving recombinant or wild-type vaccine

Test Normal ranges immunizationa immunization immunization immunizationa immunizationa immunization immunizationa immunizationa immunizationa immunizationa immunizationa immunizationa immunization immunizationa immunization immunization immunizationa immunizationa immunizationa immunization immunizationa immunizationa immunizationa immunizationa immunizationa immunizationa immunizationa immunizationa immunizational immunizationa immunizat					5	6
Normal ranges immunizationa immunizationb After immunizationa immunizationb Before immunizationa After immunizationa Before immunizationa 200-800 451 ± 48 610 ± 33 339±74 12-30 19.0 ± 3.0 17.8 ± 0.9 17.1 ± 0.6 20-60 25.2 ± 1.9 22.8 ± 1.0 28.9 ± 5.3 40-80 37.8 ± 2.3 31.8 ± 4.4 37.9 ± 3.6 200-500 194 ± 20 212 ± 21 236 ± 41 30.5-1.0 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 36, 110 662 ± 112 466 ± 119 498 ± 120			V-Wyet	h (n=4)	rV-PSA	(n=8)
Normal ranges infinitification from 33 339±74 4 200-800 451±48 610±33 339±74 4 17.1±0.6 12-30 19.0±3.0 17.8±0.9 17.1±0.6 25.2±1.9 22.8±1.0 28.9±5.3 2 20.60 37.8±2.3 31.8±4.4 37.9±3.6 3 31.8±4.4 37.9±3.6 31.8±4.4 37.9±3.6 31.8±0.03 0.8±0.05 0.8±0.05 0.8±0.05 662±112 466±119 498±120 38.00.2000		-	Before	After	Before immunization	After immunization
200-800 451±48 610±33 5.59±74 7 12-30 19.0±3.0 17.8±0.9 17.1±0.6 2 20-60 25.2±1.9 22.8±1.0 28.9±5.3 2 40-80 37.8±2.3 31.8±4.4 37.9±3.6 3 200-500 194±20 212±21 236±41 1 0.5-1.0 0.9±0.10 0.8±0.03 0.8±0.05 662±112 466±119 498±120 3		Normal ranges	HIIIIIIII Canoni		70 - 000	754 + 47
$12-30$ 19.0 ± 3.0 17.8 ± 0.9 17.1 ± 0.6 2 $20-60$ 25.2 ± 1.9 22.8 ± 1.0 28.9 ± 5.3 2 $40-80$ 37.8 ± 2.3 31.8 ± 4.4 37.9 ± 3.6 3 194 ± 20 212 ± 21 236 ± 41 194 ± 20 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 0.8 ± 0.05 0.8 ± 0.05	\ E	200-800	451±48	610 ± 33	339I /4	1 1 T 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
20-60 25.2 ± 1.9 22.8 ± 1.0 28.9 ± 5.3 2 20-60 37.8 ± 2.3 31.8 ± 4.4 37.9 ± 3.6 3 200-500 194 ± 20 212 ± 21 236 ± 41 1) 0.5-1.0 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 498 ± 120 362 ± 112 466 ± 119 498 ± 120	(1/)	10 00	100+30	17.8 ± 0.9	17.1 ± 0.6	20.5 ± 1.0
20-60 25.2 ± 1.9 22.8 ± 1.0 20.9 ± 3.5 3 $40-80$ 37.8 ± 2.3 31.8 ± 4.4 37.9 ± 3.6 3 $19/41$ $19/4 \pm 20$ 212 ± 21 236 ± 41 $19/4 \pm 20$ 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 0.8 ± 0.05 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05	g/dl)	05-71	0.5 + 0.51		20 04 5 3	258+16
40-80 37.8 ± 2.3 31.8 ± 4.4 37.9 ± 3.6 3 1g/d1) $200-500$ 194 ± 20 212 ± 21 236 ± 41 1g/d1) $0.5-1.0$ 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 1g/d1) $0.5-1.0$ 662 ± 112 466 ± 119 498 ± 120		20-60	25.2 ± 1.9	22.8 ± 1.0	C.C. T.C.07	70.04
40-8037.0 \pm 2.0212 \pm 21236 \pm 411g/dl)0.5-1.00.9 \pm 0.100.8 \pm 0.030.8 \pm 0.051g/dl)0.5-1.0662 \pm 1112466 \pm 119498 \pm 120		00 04	278+23	318+44	37.9 ± 3.6	31.9 ± 2.4
ng/dl) $0.5-1.0$ 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 $0.5-1.0$ 662 ± 1112 466 ± 119 498 ± 120	_	40-80	7.0 T 0.10		726 ± 41	104 + 13
ng/dl) $0.5-1.0$ 0.9 ± 0.10 0.8 ± 0.03 0.8 ± 0.05 in see (u/l) $500-2000$ 662 ± 112 466 ± 119 498 ± 120	_	200-500	194 ± 20	212 ± 21	14 T 057	
(u/l) 500-2000 662 ± 112 466± 119 498 ± 120			00+010	0.8 ± 0.03	0.8 ± 0.05	0.8 ± 0.02
500-2000 662 ± 112 400± 119	(mg/dl)	0.1-0.0	01:04 0:0		408 + 120	563 ± 81
	Kinase (11/		662 ± 112	400±119	071 T 071	

a 1 week prior to primary immunization b 12 weeks following primary immunization

c Alkaline phosphatase

d Blood urea nitrogen

e Alanine aminotransferase

f Aspartate aminotransferase

g Lactate dehydrogenase

Table 4 Primate IgMa Response to Inoculation with rV-PSA

		Ç			Days P	Post Immunization ^b	nization ^b		
Monkey Im	Immunogen	Dose (PFU)	pId	15	29e	43	57e	7.1	270
	V_Wweth	1×108	NDf	QN.	QN ON	QN	N ON	S	S
→ (V Wieth	1,108	C Z	Q Z	Q N	QN	QN N	Q	CZ
7 6	V Wyeth	1×108	e E	R	ON	N ON	QN N	Q.	S
ი აჭ	V-Wyeth	1×108	S	CN	CN	CIN	QN QN	CN	CN
. 4	rV.pcA	1×107	S	>40	S	20	>40	>40	QN ON
א ר	rV_pcA	1×107		>40	<u>C</u> N	CIN	20	20	CN
י כ	A DQ V.	1×107		>40	ς.	20	20	20	CN
^ သွ	rv-PSA	1×10^7	2 2	>40	5	10	>40	>40	C _N
G	rV_PSA	1×108	QN	20	S	20	10	10	QN QN
\ <u></u>	rV_PSA	1×108	QN N	20	5	40	>40	NTB	QN QN
2 =	rV-PSA	1×108	CN	>40	>40	>40	>40	>40	CZ
12c	rV-PSA	1×108	ON	>40,	20	40	20	20	CN
i									

a All monkey seras were negative for IgG to PSA at all time points; All seras were positive for IgG to vaccinia virus (>1:2000) at day 71.

b Monkeys received vaccinations on days 1, 29, and 57. Sera (1:5) was tested by ELISA. Titers were calculated using an O.D. of 0.4.

c Animal was prostatectomized.

d PI, Pre-immune.

e Animals bled before boosting.

f ND, not detectable; limit of detection was <1:5 dilution.

g NT, not tested.

the first that the control of the co

Table 5 PSA Specific Lymphoproliferative T-cell Responses of Rhesus PBMCs 270 Days Following Inoculation with rV-PSA

y-Wyeth x10 ⁸ V-Wyeth x10 ⁸ x10 ⁸ x10 ⁸ x10 ⁸ x10 ⁸ x10 ⁷ x10 ⁸ x			Doge		Ą	Antigen ^a		
V-Wyeth 1x10 ⁸ V-Wyeth 1x10 ⁸ V-Wyeth 1x10 ⁸ V-Wyeth 1x10 ⁸ V-Wyeth 1x10 ⁷ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁸	Monkey	Immunogen	(PFU)	Medium	Con A	Oval	UV-Wyeth	PSAd
V-Wyeth 1x108 V-Wyeth 1x108 V-Wyeth 1x108 V-Wyeth 1x100 rV-PSA 1x107 rV-PSA 1x107 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108		V-Wyeth	1×108	397	65701	376	24785	414
V-Wyeth 1x10 ⁸ V-Wyeth 1x10 ⁸ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁸	2b	V-Wyeth	1x108	TN	TN	Z	NT	LN
V-Wyeth 1x10 ⁸ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁷ rV-PSA 1x10 ⁸	C	V-Wyeth	1×10^8	450	84860	522	18859	413
rV-PSA 1x107 rV-PSA 1x107 rV-PSA 1x107 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108	4c	V-Wyeth	1×10^{8}	532	107840	553	16571	387
rV-PSA 1x107 rV-PSA 1x107 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108	ς,	rV-PSA	1×107	412	85276	408	6040	539
rV-PSA 1x107 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108	9	rV-PSA	1×10^7	401	96398	404	1776	3,134
rV-PSA 1x10 ⁸	7	rV-PSA	1×10^{7}	417	10806	522	10908	434
rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108	8 c	rV-PSA	1×10^{7}	1069	99216	744	15346	484
rV-PSA 1x108 rV-PSA 1x108 rV-PSA 1x108	6	rV-PSA	1×108	384	106248	386	14499	10,635
rV-PSA 1x108	10	rV-PSA	1×10^{8}	432	92263	404	19872	18,561
4V-PSA 1×108		rV-PSA	1×10^{8}	411	94055	1063	5124	16,245
	12c	rV-PSA	1×10^{8}	420	124896	392	11944	12,945

a Antigen concentrations were: Con a (2 μg/ml); Ovalbumin (100 μg/ml);
 UV-Wyeth (2x10⁷ pfu/ml); and PSA (100 μg/ml). Each value represents a mean CPM of triplicate samples. Standard deviation never exceeded 10%.

b NT, Not Tested. B-cells were not transformed for this animal.

c Animal was prostatectomized.

d Values in bold are significant when compared to their respective medium control values (p <0.001).

The state of the s

EXAMPLE II

Identification Of Potential Prostate Specific Antigen (PSA) Specific T Cell Epitopes

5

20 Inn only his hard

25

10

Since the entire amino acid sequence of human PSA is known and human class 1 HLA A2 consensus motifs have been described, studies were undertaken to identify a series of peptides that would potentially bind class 1 A2 molecules. A2 was chosen since it is the most common HLA class 1 molecule being represented in approximately 50% of North American Caucasians and 34% of African Americans. The peptide sequence of PSA was thus examined for matches to the consensus motifs for HLA A2 binding peptides. Peptides were only selected if their sequence diverged sufficiently from the PSA-related human glandular kallikrein (HGK) gene and pancreatic kallikrein antigen (PKA) sequences.

The amino acid sequence of human PSA was scanned using a predictive algorithm that combines a search for anchor residues with numerical assignments to all residues at all positions. The T2 cell binding assay was then used to determine which peptides bound human HLA A2 molecules. As can be seen in Table 6, PSA peptides 141-150, 154-163 and 146-154 scored positive in this assay (Nijman, H.W., et al., Eur. J. Immunol. 23:1215-1219, 1993). Table 7 gives the amino acid sequence of these peptides and compares them to corresponding sequences of HGK and PKA.

Table 6
PSA peptide binding assay

			PSA p	pepti	de bi	ndin	g ass	say					
	Antigen							M	1Ab	A2,	69		
5	None PSA 141 PSA 146 PSA 154	6-154 22	0.34 3.97 2.30					1	27.2	25°			
10	^a Mean c CIRA2 c	were used a hannel fluore ell line was u	scent int	ensit	.y.				A2 s	staini	ng		
15													
20			PSA pep		Tabl amin		id se	quen	ice				
	PSA	141-150	F	L	Т	Р	K	K	L	Q	С	V	
10 (10) Re	HGK		-	-	R	-	R	S	-	-	-	-	
to the training of training of the training of tra	PKA		-	S	F	-	D	D	-	-	••	-	
25 11	PSA	146-154	K	L	Q	С	V	D	L	Н	V		
Tax 2015 	HGK		S	-	-	-	-	S	-	-	L		
	PKA		D	-	-	-	-	-	-	K	l		
30	PSA	154-163	٧	entere.	S	N	D	V	С	Α	Q	٧	
	HGK		L	L	-	-	-	M	-	-	R	Α	
	PKA		l	L	Р	-	-	Ε	-	E	K	Α	

1 20 we will have all and all the subject than the subject that the subject than the subject to the subject than the subject to the subj

5

10

EXAMPLE III Establishment of Human T Cell Lines Sytolytic For Human Tumor Cells Expressing

Cytolytic For Human Tumor Cells Expressing PSA

PBMC from normal healthy donors expressing the HLA A2 class 1 allele were used in an attempt to determine if PSA specific peptides are immunogenic for humans. Peptides 141-150 and 154-163 were used in this study. The methodology used for the establishment of these cell lines involves pulsing of PBMC with peptide and IL-2 as previously described (*Tsang, K.Y., et al. JNCI, in press* and in U.S. Application Serial No. 08/396,385, the disclosure of which is herein incorporated by reference). T cell lines were able to be established from 5/6 normal donors using PSA peptide 141-150 and from 6/6 normal donors using PSA peptide 154-163. Moreover, PBMC were obtained from two prostate cancer patients. T cell lines were established from these PBMC cultures using peptide 154-163.

Some of these T cell lines have been phenotyped. As seen in Table 8, one cell line designated T-866, which was derived from pulsing with peptide 141-150, contains appreciable amounts of CD4+/CD8+ double positive cells and another cell line, designated T-1538, derived from pulsing with peptide 154-163, shows a similar phenotype.

Four of the T cell lines derived from three different individuals were then assayed for their ability to lyse human cells (Table 9). As seen in Table 9, the T cell line designated T-866, derived from peptide 141-150, was able to lyse T2 cells when pulsed with the appropriate peptide (141-150). No lysis was seen using the PSA negative human colon cancer cell line COLO-205. While 80% lysis was seen using the LNCAP PSA containing human prostate cancer cell line. When employing the NK target K562, which measures non-specific lysis due to NK cell activity, only 23% lysis was obtained. Similar results were seen employing a different T cell

25

10

line obtained from the same patient which was derived from pulsing with PSA peptide 154-163. Two additional T cell lines which were derived from peptide 154-163 were also analyzed. One was from a normal donor (T-1538) and one was from a prostate cancer patient (T-PC2). As can be seen in Table 9, employing both of these T cell lines, enhanced lysis was seen when the T2 cell line was pulsed with the 154-163 peptide and enhanced lysis was seen when employing the PSA expressing prostate specific cell line LNCAP, as compared to COLO-205 or K562. These studies demonstrate that T cell lines can be established using the peptides and protocols generated here which have the ability to lyse PSA expressing human prostate carcinoma cells.

Table 8
Flow cytometry analysis of PSA peptide specific T cells

T-cell Line	PSA Peptide	CD3	CD4	CD8	CD4/CD8	CD56
T-866	141-150	96	35	6.5	59	0
T-1538	154-163	94	5.2	32	62	0

10

15

Table 9
Cytotoxic effects of PSA peptide specific T cells

% specific release (lysis)

T-cell Line	PSA Peptide	Т2	T2+ peptide	LNCAP	K562	COLO-205
T-866	141-150	10ª	40	80	23	7
T-866	154-160	16	35	60	22	10
T-1538	154-160	10	40	29	3	10
T-PC2	154-160	15	35	35	2	8

^aPercent of ¹¹¹ In specific release 24 hour cytotoxic assay (E:T ratio, 25:1)

(SD < 2.5%)

The following is a listing of publications referred to in the foregoing specification.

5

Armbruster, D. A. Prostate-specific antigen: biochemistry, analytical methods, and clinical application. Clinical Chemistry, 39:181-195, (1993).

10

Bilhartz, D. L., Tindall, D. J., and Oesterling, J. E. Prostate-specific antigen and prostatic acid phosphatase: biomolecular and physiological characteristics. *Urology*, **38**:95-102, (1991).

Brawer, M. K., and Lange, P. H. Prostate-specific antigen and premalignant change: implications for early detection. CA Cancer Journal Clinic, 39:361-375, (1989).

15

Carter, H. B., and Coffey, D. S. The prostate: an increasing medical problem. Prostate, 16:39-48, (1990).

ment of occupants of the control of 20

Chatterjee, M. B., Foon, K. A., and Kohler, H. Idiotypic antibody immunotherapy of cancer. Cancer Immunology and Immunotherapy, **38**:75-82, (1994).

Cheever, M. A., Chen, W., Disis, M. T., and Peace, D. J. T-cell immunity to oncogenic proteins including mutated RAS and chimeric BCR-ABL. Annals of the New York Academy of Science, 690:101-112, (1993).

301

Choe, B. K., Frost, P., Morrison, M. K., and Rose, N.R. Natural killer cell activity of prostatic cancer patients. Cancer Investigations, 5:285-291, (1987).

35

Conry, R. M., Salch, M. N., Schlom, J., and LoBuglio, A. F. Breaking tolerance to carcinoembryonic antigen with a recombinant vaccinia virus vaccine in man. American Association of Cancer Research (Abstract), (1994).

Correale, P., Zaremba, S., Nieroda, C., Zhu, M. Z., Schmitz, J., Schlom, J., and Tsang, K. Y. In vitro stimulation of human cytotoxic T lymphocytes specific for peptides derived from prostate specific antigen. 9th International Congress of Immunology (Abstract), (1995).

40

Disis, M. L., Smith, J. W., Murphy, A. A., Chen, W., and Cheever, M. A. In vitro generation of human cytolytic T-cells specific for peptides from the HER-2/neu protooncogene protein. Cancer Research, 54:1071-1076, (1994).

Donovan, J. F., Lubaroff, D. M., and Williams, R. D. Immunotherapy of prostate cancer. *Problems in Urology*, 4:489-505, (1990).

Foon, K. A., Chakraborty, M., John, W., Sherratt, A., Kohler, H., and Bhattacharya-Chatterjee, M. Active immunity to the carcinoembryonic antigen (CEA) in patients treated with an anti-idiotype monoclonal antibody vaccine. *Society for Biological Therapy* (Abstract), (1994).

Gauthier, E. R., Chapdelaine, P., Tremblay, R. R., and Dube, J. Y. Characterization of rhesus monkey prostate specific antigen cDNA. *Biochimica Biophysica Acta*, **1174**:207-210, (1993).

5

10

15

20.

25

30

35

40

45

Gritz, L., Destree, A., Cormier, N., Day, E., Stallard, V., Caiazzo, T., Mazzara, G., and Panicali, D. Generation of hybrid genes and proteins by vaccinia virus-mediated recombination: application to human immunodeficiency virus type 1 env. *J. Virol.* 64:5948-5957, (1990).

Helling, F., and Livingston, P. O. Ganglioside conjugate vaccines. Immunotherapy against tumors of neuroectodermal origen. *Molecular and Chemical Neuropathology*, **21**:299-309, (1994).

Helling, F., Calves, M., Shang, Y., Oettgen, H. F., and Livingston, P. O. Construction of immunogenic GD3-conjugate vaccines. *Annals of the New York Academy of Science*, **690**:396-397, (1993).

Huang, C, L., Brassil, D., Rozzell, M., Schellhammer, P. F., and Wright, G. L. Comparison of prostate secretory protein with prostate specific antigen and prostatic acid phosphatase as a serum biomarker for diagnosis and monitoring patients with prostate carcinoma. *Prostate*, 23:201-212, (1993).

loannides, C. G., Fisk, B., Fan, D., Biddison, W. E., Wharton, J. T., and O'Brian, C. Cytotoxic T cells isolated from ovarian malignant ascites recognize a peptide derived from the HER-2/neu proto-oncogene. *Cellular Immunology*, **151**:225-234, (1993).

Irvine, K., Kantor, J., and Schlom, J. Comparison of a CEA-recombinant vaccinia virus, purified CEA, and an anti-idiotype antibody bearing the image of a CEA epitope in the treatment and prevention of CEA-expressing tumors. *Vaccine Research*, **2**:79-94, (1993).

Isaacs, J. T., Feitz, W. F., and Scheres, J. Establishment and characterization of seven Dunning rat prostatic cancer cell lines and their use in developing methods for predicting metastatic abilities of prostatic cancers. *Prostate*, **9**:261-281, (1986).

Jenkins, S., Gritz, L., Fedor, C., O'Neil, E., Cohen, L. and Panicali, D. Formation of lentivirus particles in mammalian cells infected with recombinant fowlpox virus. *AIDS Research and Human Retroviruses* 7:991-998, (1991).

5

Kantor, J., Irvine, K., Abrams, S., Kaufman, H., Dipietro, J., and Schlom, J. Antitumor activity and immune responses induced by a recombinant carcinoembryonic antigen-vaccinia virus vaccine. *Journal of the National Cancer Institute*, **84**:1084-1091, (1992a).

10

Kantor, J., Irvine, K., Abrams, S., Snoy, P., Olsen, R., Greiner, J., Kaufman, H., Eggensperger, D., and Shlom, J. Immunogenicity and safety of a recombinant vaccinia virus vaccine expressing the carcinoembryonic antigen gene in a nonhuman primate. *Cancer Research*, **52**:6917-6925, (1992b).

15

Karr, J. F., Kantor, J. A., Hand, P. H., Eggensperger, D. L., and Schlom, J. Conservation of the prostate specific antigen (PSA) gene in primates and the expression of recombinant human PSA in a transfected murine cell line. *Cancer Research*: Submitted for Publication, (1995).

20 min min libra

Kaufman, H., Schlom, J., and Kantor, J. A recombinant vaccinia virus expressing human carcinoembryonic antigen (CEA). *International Journal of Cancer*, **48**:900-907, (1991).

Kleer, E., and Oesterling, J. E. PSA and staging of localized prostate cancer. *Urologic Clinics of North America*, **20**:695-704, (1993).

Lilja, H. Structure, function, and regulation of the enzyme activity of prostate-specific antigen. *World Journal of Urology*, **11**:188-191, (1993).

35

30

Livingston, P. O., Calves, M. J., Helling, F., Zollinger, W. D., Blake, M. S., and Lowell, G. H. GD3/proteosome vaccines induce consistent IgM

antibodies against the ganglioside GD3. Vaccine, 12:1199-1204, (1993).

Lundwall, A., and Lilja, H. Molecular cloning of human prostate specific antigen cDNA. *FEBS Letters*, **214**:317-322, (1987).

40

McEntee, M., Isaacs, W., and Smith, C. Adenocarcinoma of the canine prostate: immunohistochemical examination for secretory antigens. *Prostate*, **11**:163-170, (1987).

45

Moss, B. Generation of recombinant vaccinia viruses. *Current Protocols in Molecular Biology*, 2:16.15.1-16.18.9, (1993).

- Oesterling, J. E. Prostate specific antigen: a critical assessment of the most useful tumor marker for adenocarcinoma of the prostate. *Journal of Urology*, **145**:907-923, (1991).
- Panicali, D., Grzelecki, A. and Huang, C. Vaccinia virus vectors utilizing the β-galactosidase assay for rapid selection of recombinant viruses and measurement of gene expression. *Gene* 47:193-199, (1986).

15

Hard delay. The second of the

20

The House with

25

30

35

40

- Paoletti, E., Tartaglia, J., and Cox, W. I. Immunotherapeutic stratagies for cancer using poxvirus vectors. *Annals of the New York Academy of Sciences*, **690**:292-300, (1993).
- Peace, D. J., Xue, B., Sosman, J. A., and Zhang, Y. In vitro immunization of human cytotoxic T lymphocytes specific for peptides derived from prostate specific antigen. *Cancer Vaccines: Structural Basis for Vaccine Development* (Abstract), (1994).
- Powrie, F., and Coffman, R. L. Cytokine regulation of T-cell function: potential for therapeutic intervention. *Immunology Today*, **14**:270-274, (1993).
- Ravindranath, M. H., Brazeau, S. M., and Morton, D. L. Efficacy of tumor cell vaccine after incorporating monophosphoryl A (MPL) in tumor cell membranes containing tumor associated ganglioside. *Experimentia*, **50**:648-653, (1994).
- Ritter, G., Boosfeld, E., Adluri, R., Calves, M., Oettgen, H. F., Old, L. J., and Livingston, P. Antibody response to immunization with ganglioside GD3 and GD3 congeners (lactones, amide, and ganglisidol) in patients with malignant melanoma. *International Journal of Cancer*, **48**:379-385, (1991).
- Schellhammer, P. F., and Wright, G. L. Biomolecular and clinical characteristics of PSA and other candidate prostate tumor markers. *Urologic Clinics of North America*, **20**:597-606, (1993).
- Schlom, J., Kantor, J., Abrams, S., Tsang, K. Y., Panicali, D., and Hamilton, J. M. Strategies for the development of recombinant vaccines for the immunotherapy of breast cancer. *Breast Cancer Research and Treatment*, In Press.
- Schroder, F. H. Experimental Models in the study of prostate cancer. Prostate Cancer. In: *International Perspectives in Urology.*, **3**:343-377, (1982).

10

15

Tsang, K. Y., Nieroda, C. A., De Filippi, R., Chung, Y. K., Yamaue, H., Greiner, J. W., and Schlom, J. Induction of human cytotoxic T cell lines directed against point-mutated p21 Ras-derived synthetic peptides. *Vaccine Research*, **3**:183-193, (1994).

Wakui, S., Furusato, M., Nomura, Y., Asari, M., and Kano, Y. Lectin histochemical study of the prostate gland of the rhesus monkey (Macaca mulatta). *Journal of Anatomy*, **181**:127-131, (1992).

Wang, M. C., Kuriyama, M., Papsidero, L. D., Loor, R. M., Valenzuela, L. A., Murphy, G. P., and Chu, T. M. Prostate antigen of human cancer patients. *Methods in Cancer Research*, **19**:179-197, (1982).

Wang, M. C., Valenzuela, L. A., Murphy, G. P., and Chu, T. M. Purification of a human prostate specific antigen. *Investigations in Urology*, **17**:159-163, (1979).

Zietman, A. L., Shipley, W. L., and Willett, C. G. Residual disease after radical surgery or radiation therapy for prostate cancer. Clinical significance and therapeutic implications. *Cancer*, 71:959-969, (1993).

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: SCHLOM, JEFFREY PANICALI, DENNIS
 - (ii) TITLE OF INVENTION: GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC ANTIGEN (PSA)
 - (iii) NUMBER OF SEQUENCES: 2
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: SEWALL P. BRONSTEIN; DIKE, BRONSTEIN, ROBERTS & CUSHMAN
 - (B) STREET: 130 WATER STREET
 - (C) CITY: BOSTON
 - (D) STATE: MASSACHUSETTS
 - (E) COUNTRY: US
 - (F) ZIP: 02129
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: Patentln Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: RESNICK, DAVID S.
 - (B) REGISTRATION NUMBER: 34,235
 - (C) REFERENCE/DOCKET NUMBER: 44981
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (617) 523-3400
 - (B) TELEFAX: (617) 523-6440
 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TCTAGAAGCC CCAAGCTTAC CACCTGCA

28

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 41 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

TCTAGATCAG GGGTTGGCCA CGATGGTGTC CTTGATCCAC T

This invention has been described in detail including the preferred embodiments thereof. However, it will be appreciated that those skilled in the art, upon consideration of this disclosure, may make modifications and improvements thereon without departing from the spirit and scope of the invention as set forth in the claims.

What is claimed is:

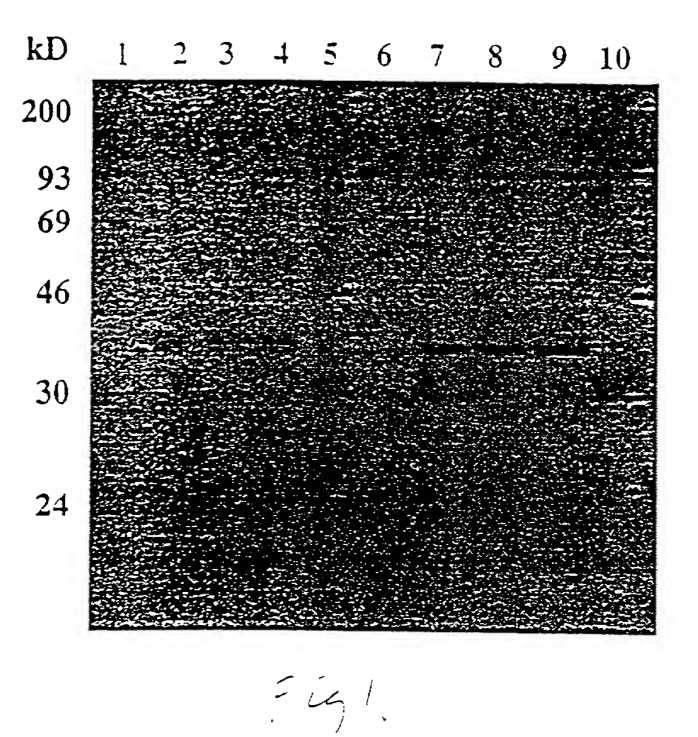
- 1. A method for generating an immune response to prostate-specific antigen (PSA) comprising, introducing a sufficient amount of a first pox virus vector to a host to stimulate an immune response, wherein the pox virus vector has at least one insertion site containing a DNA segment encoding PSA operably linked to a promoter capable of expression in the host.
- 2. The method of claim 1, further comprising at at least one periodic interval after introduction of the first pox virus vector contacting the host with additional PSA or a cytotoxic T-cell eliciting epitope thereof.
- 3. The method of claim 2, wherein the host is contacted with the additional PSA by introducing a second pox virus vector to the host having at least one insertion site containing a DNA segment encoding the PSA operably linked to a promoter capable of expression in the host.
- 4. A method for generating an immune response to prostatespecific antigen (PSA) in a host, comprising:
- a. contacting the host with a sufficient amount of PSA or a cytotoxic T-cell eliciting epitope thereof; and
- b. at least one periodic interval thereafter contacting the host with additional PSA or a cytotoxic T-cell eliciting epitope thereof.
- 5. The method of claim 4, wherein the host is contacted with the additional PSA by introducing a pox virus vector to the host having at least one insertion site containing a DNA segment encoding PSA or a cytotoxic T-cell eliciting epitope thereof operably linked to a promoter capable of expression in the host.

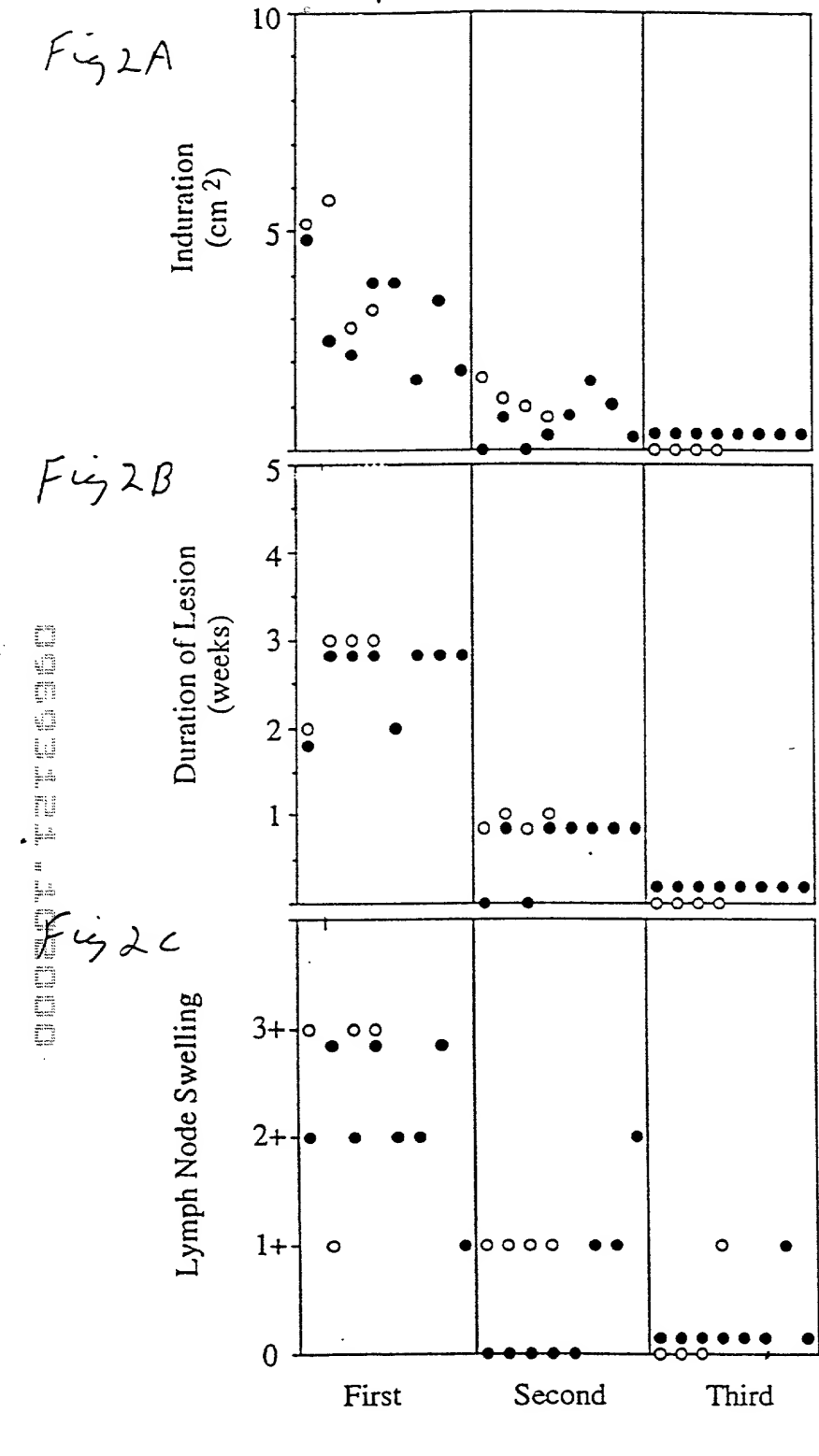
- 6. The method of claim 1 or 5, wherein the pox virus is selected from the group of pox viruses consisting of suipox, avipox, capripox and orthopox virus.
- 7. The method of claim 6, wherein the orthopox virus is vaccinia.
- 8. The method of claim 7, wherein the avipox is fowlpox, canary pox and pigeon pox.
 - 9. The method of claim 8, wherein the suipox is swinepox.
- 10. The method of claim 3, wherein the first pox virus vector is vaccinia and the second pox virus vector is selected from the group of pox viruses consisting of suipox, avipox, capripox and orthopox virus.
- 11. The method of claim 2 or 4, wherein the PSA or T-cell eliciting epitope is formulated with an adjuvant or is in a liposomal formulation.
- 12. The method of claim 11, wherein the adjuvant is selected from the group consisting of RIBI Detox, QS21 and incomplete Freund's adjuvant.
- 13. A method for generating an immune response to PSA comprising contacting a host with a cytotoxic T-cell eliciting epitope of PSA.

- 14. The method of claim 13, wherein the T-cell eliciting epitope is formulated with an adjuvant or is in a liposomal formulation.
- 15. The method of claim 12, wherein the adjuvant is selected from the group consisting of RIBI Detox, QS21 and incomplete Freund's adjuvant.
- 16. A pharmaceutical composition comprising a pox virus vector having at least one insertion site containing a DNA reagent encoding PSA operably linked to a promoter and a pharmaceutical carrier.

ABSTRACT

We have discovered that by using a recombinant viral vector, preferably a pox virus vector having at least one insertion site containing a DNA segment encoding prostate-specific antigen (PSA), operably linked to a promoter capable of expression in the host, a specific humoral and cellular immune response to PSA can be generated. The method preferably comprises introducing a sufficient amount of the recombinant pox virus vector into a host to stimulate the immune response, and contacting the host with additional PSA at periodic intervals thereafter. The additional PSA may be added by using a second pox virus vector from a different pox genus. In another embodiment, additional PSA can be added by contacting the host with PSA by a variety of other methods, including in one preferred embodiment adding PSA. The PSA may be formulated with an adjuvant or in a liposomal formulation.





Inoculation

CIKE, BRONSTEIN, ROBERTS & CUSHMAN 130 Water Street Boston

DECLARATION Original Application

ATTORNEY'S DOCKET NO (II Any) AND POWER OF ATTORNEY 45394 Massachusetts 02109 As a below named inventor, I hereby declare that My residence, post office address and citizenship are as stated below next to my name, I believe I am the original, first and sole inventor (if only one name is listed at 201 below) or an original, first and joint inventor (if at 201-203 below) of the subject matter which is claimed and for which a patent is sought on the invention entitled GENERATION OF IMMUNE RESPONSES TO PROSTATE SPECIFIC ANTIGEN (PSA) which is described and claimed in: xx the specification in application Serial No. 08/500,306 tiled ...July..10,..1995. the attached specification or (for declaration not accompanying application) if applicable I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by ans amendment referred to above Facknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a). I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed

		YES	
		1	NO
		YES	NO
PRIOR TO THE F	ILING DATE OF THIS APPLICA	ATION	
	States application	States application(s) listed below and, insofar	States application(s) listed below and, insofar as the subjet States application in the manner provided by the first se material information as defined in Title 37, Code of

of this application: (Application Serial No.) (Filing Date) (Status) (Patented, pending, abandoned)

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) with full powers of association, substitution and revocation to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (list name and registration number)

Sewali P. Bronstein (Reg. No. 16,919) Ernest V. Linek (Reg. No. 29,822) Peter J. Manus (Reg. No. 26,766) Brian L. Michaelis (Reg. No. 34,221) Donald Brown (Reg. No. 20,845) Linda M. Buckley (Reg. No. 31,003) David S. Resnick (Reg. No. 34,235) Cara Z. Lowen (Reg. No. 38,227) David G. Conlin (Reg. No. 27,026) Ronald I. Eisenstein (Reg. No. 30,628) Peter F. Corless (Reg. No. 33,860) John L. Welch (Reg. No. 28,129) George W. Neuner (Reg. No. 26,964) Henry D. Pahl, Jr. (Reg. No. 20,438) Kevin J. Fournier (Reg. No. 34,333) Milton Mc.K. Oliver (Reg. No. 28,333)

SEND CORRESPONDENCE TO: DIRECT TELEPHONE CALLS TO: Sewall P. Bronstein, Esq. (name and telephone number) Dike, Bronstein, Roberts & Cushman David S. Resnick 130 Water Street Boston, Massachusetts 02109 617/523-3400

	FULL NAME	LAST NAME	FIRST NAME	MIDDLE NAME	
	OF INVENTOR	SCHLOM	Jeffrey		
23	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZE	NSHIP
2	CITIZENSHIP	Potomac	Maryland	U.S.A.	
	DOCT OFFICE	POST OFFICE ADDRESS	CITY	STATE OR COUNTRY	ZIP CODE
	POST OFFICE ADDRESS	10301 Sorrel Ave.	Potomac	Maryland	20854
	FULL NAME	LAST NAME	FIRST NAME	MIDDLE NAME	***************************************
	OF INVENTOR	PANICALI	Dennis	I.a	
2	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZE	NSHIP
202	CITIZENSHIP	Acton	Massachusetts	U.S.A.	
	DOCT OFFICE	POST OFFICE ADDRESS	CITY	STATE OR COUNTRY	ZIP CODE
	POST OFFICE ADDRESS	114 Nonset Path	Acton	Massachusetts	01720
	FULL NAME OF INVENTOR	LAST NAME	FIRST NAME	MIDDLE NAME	
203	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZEI	NSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS	CITY	STATE OR COUNTRY	ZIP CODE

I hereby further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

SIGNATURE OF INVENTOR 201	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
DATE	10-25-95	DATE

U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE The state of the s

DECLARATION AND POWER OF ATTORNEY Original Application

ATTORNEY'S DOCKET NO (II Any) 45394

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed at 20) below) or an original, first and joint inventor (if plural names are based

at 		I the subject matter which is ON OF IMMUNE RESP								
_	nich is described a	estication of XXI		ion in application			,306	nledJu1	y10,	1995
		, ,	and was amo	ended on						• • • • • • •
	amendment tele	ie duty to disclose information					n, mch	•		
	listed below and	26(a). oreign priority benefits under have also identified below at high priority is claimed	Title 35, Uny foreign ap	nited States Code, pplication for pate	, \$119 of any intentor	foreign ap s certifica	plicatio ite havii	n(s) for patent or inven ng a filing date belore	itor's certi that of the	ficate :
		OREIGN APPLICATION(S), IF	ANY, FILED	WITHIN 12 MONT	HS PRIOR TO	THE FILIN	G DATE	OF THIS APPLICATION		
		COUNTRY			APPLICATIO			DATE OF FILING (day, month, year)	PRIO CLAIMED	
									YES	ОМ
									YES	МО
	ALL F	OREIGN APPLICATIONS, IF A	NY, FILED N	AORE THAN 12 M	ONTHS PRIOR	TO THE F	ILING (DATE OF THIS APPLICA	TION	
										
						·····	_	· · · · · · · · · · · · · · · · · · ·		
	matter of each of Daragraph of Tit	ne benefit under Title 15, Unit the claims of this application le 35, United States Code, §1 ons, §1.56(a) which occurred on.	on is not disc 112, Lackno	closed in the prio wledge the duty t	r United States o disclose mate	- applicati rial intor	on m tl mation	se manner provided by as defined in Title 37,	the first Cod e of	
11	Application Serial	No)		(1 ding 1)	ate)		(Status) (Patented, pending, :	shandoned	i)
							·			
	revocation to pro registration num. Sewall P. Bronstein Donald Brown David G. Conlin	(Reg. No. 16,919) Ernest V (Reg. No. 20,845) Linda M (Reg. No. 27,026) Ronald I	ransact all b '. Linek (F '. Buckley (F '. Eisenstein (F	reby appoint the usiness in the Pat leg. No. 29,822) leg. No. 31,003) leg. No. 30,628) leg. No. 20,438)	following attorent and Trader Peter J. Manus David S. Resnic Peter F. Corless Kevin J. Fourni	(Reg. ? k tHeg. ? i (Reg. ?	th full pice conn No. 26,766 No. 34,235 No. 33,860 No. 34,333	Brian L. Michaelis Cara Z. Lowen John L. Welch	ine and (Reg. No. (Reg. No. (Reg. No.	. 34,221) . 38,227) . 28,129)
	SEND CORRE	SPONDENCE TO: Bronstein, Esq.						EPHONE CALLS To telephone number)	O :	
		, Roberts & Cushman t				,	ıs.	Resnick		
								MIDDLE NAME		
	FULL NAME	SCHLOM		Jeffrey				MIDDLE NAME		
-	OF INVENTOR	CITY		STATE OR FORE	IGN COUNTRY	·		COUNTRY OF CITIZEN	ISHIP	
201	RESIDENCE & CITIZENSHIP	Potomac		Maryland				U.S.A.	120.00	
	POST OFFICE	POST OFFICE ADDRESS	-	CITY			1	TE OR COUNTRY	21P CC	
-	ADDRESS	10301 Sorrel Av	e.	Potomac FIRST NAME			_ Fla	MIDDLE NAME	12003	<u></u>
	FULL NAME OF INVENTOR	PANICALI		Dennis				_L		
202	RESIDENCE &	CITY		STATE OR FORE		•		COUNTRY OF CITIZER	ISHIP	
~	CITIZENSHIP	Acton POST, OFFICE ADDRESS		Massachu	setts		SIA	U.S.A.	ZIP CC	ODE
	POST OFFICE	114 Nonset Path		Acton				ssachusetts	0172	20
\neg	FULL NAME	LAST NAME		FIRST NAME				MIDDLE NAME		
	OF INVENTOR	CITY		STATE OR FORE	IGN COUNTRY			COUNTRY OF CITIZEN	SHIP	
33	RESIDENCE &	Citt		JANE ON FONE						
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY			STA	TE OR COUNTRY	ZIP CO	DE
	are believed in h	declare that all statements may true; and further that these is or imprisonment, or both,	statements:	were madê wilh t	he knowledge I	hat willlu	il laise i	Mateurine and the nee	W IIIAUE 4	411

may jeopardize the validity of the application or any patent issued thereon.

SIGNATURE OF INVENTOR 291	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
00EN 11/20/95	DATE	DATE

U.S. DEPARTMENT OF COMMERCE-PATENT AND TRADEMARK OFFICE

Practitioner's Docket No. 700953-45394-CPA-2



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

Schlom et al.

Application No.:

Continuation of 08/500,306

Filed:

July 10, 1995

For:

GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC

ANTIGEN (PSA)

Assistant Commissioner for Patents Washington, D.C. 20231

CHANGE OF ATTORNEY'S ADDRESS IN APPLICATION

Please send all correspondence for this application as follows:

Ronald I. Eisenstein NIXON PEABODY LLP 101 Federal Street Boston, MA 02110

Please direct telephone calls to:

(617) 345-6054

Reg. No. 34,235

Tel. No.: (617) 345-6057

SIGNATURE OF PRACTITIONER

David Resnick

NIXON PEABODY LLP

101 Federal Street Boston, MA 02110

CERTIFICATE OF MAILING/TRANSMISSION (37 C.F.R. 1.8(a))

I hereby certify that, on the date shown below, this correspondence is being:

MAILING

deposited with the United States Postal Service with sufficient postage as first class mail in an envelope addressed to the Assistant Commissioner for Patents, Washington, D.C. 20231.

Date: 10/20/00

FACSIMILE

transmitted by facsimile to the Patent and Trademark Office.

Datricia l'urner Signature

Patricia Turner

(Change of Attorney's Address in Application, page 1 of 1)

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: SCHLOM, JEFFREY PANICALI, DENNIS
- (ii) TITLE OF INVENTION: GENERATION OF IMMUNE RESPONSES TO PROSTATE-SPECIFIC ANTIGEN (PSA)
- (iii) NUMBER OF SEQUENCES: 2
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: SEWALL P. BRONSTEIN; DIKE, BRONSTEIN, ROBERTS & CUSHMAN
 - (B) STREET: 130 WATER STREET
 - (C) CITY: BOSTON
 - (D) STATE: MASSACHUSETTS
 - (E) COUNTRY: US
 - (F) ZIP: 02129
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: RESNICK, DAVID S.
 - (B) REGISTRATION NUMBER: 34,235
 - (C) REFERENCE/DOCKET NUMBER: 44981
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (617) 523-3400
 - (B) TELEFAX: (617) 523-6440
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown

The state of the s

100 mm
Man de la companya de
7 1 1 1 1 1
ŧ
10 m
Hann,
11 (12) 11 (12)
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

TCTAGAAGCC CCAAGCTTAC CACCTGCA	28
(2) INFORMATION FOR SEQ ID NO:2:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: unknown 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
TCTAGATCAG GGGTTGGCCA CGATGGTGTC CTTGATCCAC T	41

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1: